Ath....ey Docket No.: OPHD-02304

MULTIVALENT VACCINE FOR CLOSTRIDIUM BOTULINUM NEUROTOXIN

This application is a Continuation-In-Part of copending Application Serial No. 08/405,496, filed March 16, 1995.

FIELD OF THE INVENTION

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The present invention relates to the isolation of polypeptides derived from Clostridium botulinum neurotoxins and the use thereof as immunogens for the production of vaccines, including multivalent vaccines, and antitoxins.

BACKGROUND OF THE INVENTION

The genus Clostridium is comprised of gram-positive, anaerobic, spore-forming bacilli. The natural habitat of these organisms is the environment and the intestinal tracts of humans and other animals. Indeed, clostridia are ubiquitous; they are commonly found in soil, dust, sewage, marine sediments, decaying vegetation, and mud. [See e.g., P.H.A. Sneath et al., "Clostridium," Bergey's Manual® of Systematic Bacteriology, Vol. 2, pp. 1141-1200, Williams & Wilkins (1986).] Despite the identification of approximately 100 species of Clostridium, only a small number have been recognized as etiologic agents of medical and veterinary importance.

Nonetheless, these species are associated with very serious diseases, including botulism, tetanus, anaerobic cellulitis, gas gangrene, bacteremia, pseudomembranous colitis, and clostridial gastroenteritis. Table I lists some of the species of medical and veterinary importance and the diseases with which they are associated. As virtually all of these species have been isolated from fecal samples of apparently healthy persons, some of these isolates may be transient, rather than permanent residents of the colonic flora.

TABLE 1

Clostridium Species Of Medical And Veterinary Importance*

	Species	Disease
	C. aminovalericum	Bacteriuria (pregnant women)
5	C. argentinense	Infected wounds; Bacteremia; Botulism; Infections of amniotic fluid
	C. baratii	Infected war wounds; Peritonitis; Infectious processes of the eye, ear and prostate
	C. beijerinckikii	Infected wounds
10	C. bifermentans	Infected wounds; Abscesses; Gas Gangrene; Bacteremia
	C. botulinum	Food poisoning; Botulism (wound, food, infant)
	C. butyricum	Urinary tract, lower respiratory tract, pleural cavity, and abdominal infections; Infected wounds; Abscesses; Bacteremia
	C. cadaveris	Abscesses; Infected wounds
	C. carnis	Soft tissue infections; Bacteremia
	C. chawoei	Blackleg
15	C. clostridioforme	Abdominal, cervical, scrotal, pleural, and other infections; Septicemia; Peritonitis; Appendicitis
	C. cochlearium	Isolated from human disease processes, but role in disease unknown.
	C. difficile	Antimicrobial-associated diarrhea; Pseudomembranous enterocolitis; Bacteremia; Pyogenic infections
20	C. fallax	Soft tissue infections
	C. ghnoii	Soft tissue infections
	C. glycolicum	Wound infections; Abscesses; Peritonitis
	C. hastiforme	Infected war wounds; Bacteremia; Abscesses
	C. histolyticum	Infected war wounds; Gas gangrene; Gingival plaque isolate
	C. indolis	Gastrointestinal tract infections
	C. innocuum	Gastrointestinal tract infections; Empyema
25	C. irregulare	Penile lesions
	C. leptum	Isolated from human disease processes, but role in disease unknown.
	C. limosum	Bacteremia; Peritonitis; Pulmonary infections

TABLE 1

Clostridium Species Of Medical And Veterinary Importance*

Species	Disease
C. malenominatum	Various infectious processes
C. novyi	Infected wounds; Gas gangrene; Blackleg, Big head (ovine); Redwater disease (bovine)
C. oroticum	Urinary tract infections; Rectal abscesses
C. paraputrificum	Bacteremia; Peritonitis; Infected wounds; Appendicitis
C. perfringens	Gas gangrene; Anaerobic cellulitis; Intra-abdominal abscesses; Soft tissue infections; Food poisoning; Necrotizing pneumonia; Empyema; Meningitis; Bacteremia; Uterine Infections; Enteritis necrotans; Lamb dysentery; Struck; Ovine Enterotoxemia;
C. putrefaciens	Bacteriuria (Pregnant women with bacteremia)
C. putrificum	Abscesses; Infected wounds; Bacteremia
C. ramosum	Infections of the abdominal cavity, genital tract, lung, and biliar tract; Bacteremia
C. sartagoforme	Isolated from human disease processes, but role in disease unknown.
C. septicum	Gas gangrene; Bacteremia; Suppurative infections; Necrotizing enterocolitis; Braxy
C. sordellii	Gas gangrene; Wound infections; Penile lesions; Bacteremia; Abscesses; Abdominal and vaginal infections
C. sphenoides	Appendicitis; Bacteremia; Bone and soft tissue infections; Intraperitoneal infections; Infected war wounds; Visceral gas gangrene; Renal abscesses
C. sporogenes	Gas gangrene; Bacteremia; Endocarditis; central nervous system and pleuropulmonary infections; Penile lesions; Infected war wounds; Other pyogenic infections
C. subterminale	Bacteremia; Empyema; Biliary tract, soft tissue and bone infections
C. symbiosum	Liver abscesses; Bacteremia; Infections resulting due to bowel flora
C. tertium	Gas gangrene; Appendicitis; Brain abscesses; Intestinal tract and soft tissue infections; Infected war wounds; Periodontitis; Bacteremia

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TABLE 1

Clostridium Species Of Medical And Veterinary Importance*

Species	Disease
C. tetani	Tetanus; Infected gums and teeth; Corneal ulcerations; Mastoid and middle ear infections; Intraperitoneal infections; Tetanus neonatorum; Postpartum uterine infections; Soft tissue infections, especially related to trauma (including abrasions and lacerations); Infections related to use of contaminated needles
C. thermosaccharolyticum	Isolated from human disease processes, but role in disease unknown.

Compiled from P.G. Engelkirk et al. "Classification", Principles and Practice of Clinical Anaerobic Bacteriology, pp. 22-23, Star Publishing Co., Belmont, CA (1992); J. Stephen and R.A. Petrowski, "Toxins Which Traverse Membranes and Deregulate Cells," in Bacterial Toxins, 2d ed., pp. 66-67, American Society for Microbiology (1986); R. Berkow and A.J. Fletcher (eds.), "Bacterial Diseases," Merck Manual of Diagnosis and Therapy, 16th ed., pp. 116-126, Merck Research Laboratories, Rahway, N.J. (1992); and O.H. Sigmund and C.M. Fraser (eds.), "Clostridial Infections," Merck Veterinary Manual, 5th ed., pp. 396-409, Merck & Co., Rahway, N.J. (1979).

In most cases, the pathogenicity of these organisms is related to the release of powerful exotoxins or highly destructive enzymes. Indeed, several species of the genus *Clostridium* produce toxins and other enzymes of great medical and veterinary significance. [C.L. Hatheway, Clin. Microbiol. Rev. 3:66-98 (1990).]

Perhaps because of their significance for human and veterinary medicine, much research has been conducted on these toxins, in particular those of *C. botulinum* and *C. difficile*.

C. botulinum

Several strains of Clostridium botulinum produce toxins of significance to human and animal health. [C.L. Hatheway, Clin. Microbiol. Rev. 3:66-98 (1990)] The effects of these toxins range from diarrheal diseases that can cause destruction of the colon, to paralytic effects that can cause death. Particularly at risk for developing clostridial diseases are neonates and humans and animals in poor health (e.g., those suffering from diseases associated with old age or immunodeficiency diseases).

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Clostridium botulinum produces the most poisonous biological toxin known. The lethal human dose is a mere 10-9 mg/kg bodyweight for toxin in the bloodstream. Botulinal toxin blocks nerve transmission to the muscles, resulting in flaccid paralysis. When the toxin reaches airway and respiratory muscles, it results in respiratory failure that can cause death. [S. Arnon, J. Infect. Dis. 154:201-206 (1986)]

C. botulinum spores are carried by dust and are found on vegetables taken from the soil, on fresh fruits, and on agricultural products such as honey. Under conditions favorable to the organism, the spores germinate to vegetative cells which produces toxin. [S. Arnon, Ann. Rev. Med. 31:541 (1980)]

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Botulism disease may be grouped into four types, based on the method of introduction of toxin into the bloodstream. Food-borne botulism results from ingesting improperly preserved and inadequately heated food that contains botulinal toxin. There were 355 cases of food-borne botulism in the United States between 1976 and 1984. [K.L. MacDonald et al., Am. J. Epidemiol. 124:794 (1986).] The death rate due to botulinal toxin is 12% and can be higher in particular risk groups. [C.O. Tacket et al., Am. J. Med. 76:794 (1984).] Wound-induced botulism results from C. botulinum penetrating traumatized tissue and producing toxin that is absorbed into the bloodstream. Since 1950, thirty cases of wound botulism have been reported. [M.N. Swartz, "Anaerobic Spore-Forming Bacilli: The Clostridia," pp. 633-646, in B.D. Davis et al., (eds.), Microbiology, 4th edition, J.B. Lippincott Co. (1990).] Inhalation botulism results when the toxin is inhaled. Inhalation botulism has been reported as the result of accidental exposure in the laboratory [E. Holzer, Med. Klin. 41:1735 (1962)] and could arise if the toxin is used as an agent of biological warfare [D.R. Franz et al., in Botulinum and Tetanus Neurotoxins, B.R. DasGupta, ed., Plenum Press, New York (1993), pp. 473-476]. Infectious infant botulism results from C. botulinum colonization of the infant intestine with production of toxin and its absorption into the bloodstream. It is likely that the bacterium gains entry when

spores are ingested and subsequently germinate. [S. Amon, J. Infect. Dis. 154:201 (1986).] There have been 500 cases reported since it was first recognized in 1976. [M.N. Swartz, supra.]

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Infant botulism strikes infants who are three weeks to eleven months old (greater than 90% of the cases are infants less than six months). [S. Arnon, J. Infect. Dis. 154:201 (1986).] It is believed that infants are susceptible, due, in large part, to the absence of the full adult complement of intestinal microflora. The benign microflora present in the adult intestine provide an acidic environment that is not favorable to colonization by *C. botulinum*. Infants begin life with a sterile intestine which is gradually colonized by microflora. Because of the limited microflora present in early infancy, the intestinal environment is not as acidic, allowing for *C. botulinum* spore germination, growth, and toxin production. In this regard, some adults who have undergone antibiotic therapy which alters intestinal microflora become more susceptible to botulism.

An additional factor accounting for infant susceptibility to infectious botulism is the immaturity of the infant immune system. The mature immune system is sensitized to bacterial antigens and produces protective antibodies. Secretory IgA produced in the adult intestine has the ability to agglutinate vegetative cells of C. botulinum. [S. Arnon, J. Infect. Dis. 154:201 (1986).] Secretory IgA may also act by preventing intestinal bacteria and their products from crossing the cells of the intestine. [S. Arnon, Epidemiol. Rev. 3:45 (1981).] The infant immune system is not primed to do this.

Clinical symptoms of infant botulism range from mild paralysis, to moderate and severe paralysis requiring hospitalization, to fulminant paralysis, leading to sudden death. [S. Amon, Epidemiol. Rev. 3:45 (1981).]

The chief therapy for severe infant botulism is ventilatory assistance using a mechanical respirator and concurrent elimination of toxin and bacteria using cathartics,

enemas, and gastric lavage. There were 68 hospitalizations in California for infant botulism in a single year with a total cost of over \$4 million for treatment.

[T.L. Frankovich and S. Arnon, West. J. Med. 154:103 (1991).]

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Different strains of Clostridium botulinum each produce antigenically distinct toxin designated by the letters A-G. Serotype A toxin has been implicated in 26% of the cases of food botulism; types B, E and F have also been implicated in a smaller percentage of the food botulism cases [H. Sugiyama, Microbiol. Rev. 44:419 (1980)]. Wound botulism has been reportedly caused by only types A or B toxins [H. Sugiyama, supra]. Nearly all cases of infant botulism have been caused by bacteria producing either type A or type B toxin. (Exceptionally, one New Mexico case was caused by Clostridium botulinum producing type F toxin and another by Clostridium botulinum producing a type B-type F hybrid.) [S. Arnon, Epidemiol. Rev. 3:45 (1981).] Type C toxin affects waterfowl, cattle, horses and mink. Type D toxin affects cattle, and type E toxin affects both humans and birds.

A trivalent antitoxin derived from horse plasma is commercially available from Connaught Industries Ltd. as a therapy for toxin types A, B, and E. However, the antitoxin has several disadvantages. First, extremely large dosages must be injected intravenously and/or intramuscularly. Second, the antitoxin has serious side effects such as acute anaphylaxis which can lead to death, and serum sickness. Finally, the efficacy of the antitoxin is uncertain and the treatment is costly. [C.O. Tacket et al., Am. J. Med. 76:794 (1984).]

A heptavalent equine botulinal antitoxin which uses only the F(ab')2 portion of the antibody molecule has been tested by the United States Military. [M. Balady, USAMRDC Newsletter, p. 6 (1991).] This was raised against impure toxoids in those large animals and is not a high titer preparation.

A pentavalent human antitoxin has been collected from immunized human subjects for use as a treatment for infant botulism. The supply of this antitoxin is

limited and cannot be expected to meet the needs of all individuals stricken with botulism disease. In addition, collection of human sera must involve screening out HIV and other potentially serious human pathogens. [P.J. Schwarz and S.S. Arnon, Western J. Med. 156:197 (1992).]

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Infant botulism has been implicated as the cause of mortality in some cases of Sudden Infant Death Syndrome (SIDS, also known as crib death). SIDS is officially recognized as infant death that is sudden and unexpected and that remained unexplained despite complete post-mortem examination. The link of SIDS to infant botulism came when fecal or blood specimens taken at autopsy from SIDS infants were found to contain *C. botulinum* organisms and/or toxin in 3-4% of cases analyzed. [D.R. Peterson *et al.*, Rev. Infect. Dis. 1:630 (1979).] In contrast, only 1 of 160 healthy infants (0.6%) had *C. botulinum* organisms in the feces and no botulinal toxin. (S. Arnon *et al.*, Lancet, pp. 1273-76, June 17, 1978.)

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In developed countries, SIDS is the number one cause of death in children between one month and one year old. (S. Arnon et al., Lancet, pp. 1273-77, June 17, 1978.) More children die from SIDS in the first year than from any other single cause of death in the first fourteen years of life. In the United States, there are 8,000-10,000 SIDS victims annually. *Id*.

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What is needed is an effective therapy against infant botulism that is free of dangerous side effects, is available in large supply at a reasonable price, and can be safely and gently delivered so that prophylactic application to infants is feasible.

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Immunization of subjects with toxin preparations has been done in an attempt to induce immunity against botulinal toxins. A C. botulinum vaccine comprising chemically inactivated (i.e., formaldehyde-treated) type A, B, C, D and E toxin is commercially available for human usage. However, this vaccine preparation has several disadvantages. First, the efficacy of this vaccine is variable (in particular, only 78% of recipients produce protective levels of anti-type B antibodies following

administration of the primary series). Second, immunization is painful (deep subcutaneous inoculation is required for administration), with adverse reactions being common (moderate to severe local reactions occur in approximately 6% of recipients upon initial injection; this number rises to approximately 11% of individuals who receive booster injections) [Informational Brochure for the Pentavalent (ABCDE) Botulinum Toxoid, Centers for Disease Control]. Third, preparation of the vaccine is dangerous as active toxin must be handled by laboratory workers.

What is needed are safe and effective vaccine preparations for administration to those at risk of exposure to *C. botulinum* toxins.

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C. difficile

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C. difficile, an organism which gained its name due to difficulties encountered in its isolation, has recently been proven to be an etiologic agent of diarrheal disease. (Sneath et al., p. 1165.). C. difficile is present in the gastrointestinal tract of approximately 3% of healthy adults, and 10-30% of neonates without adverse effect (Swartz, at p. 644); by other estimates, C. difficile is a part of the normal gastrointestinal flora of 2-10% of humans. [G.F. Brooks et al., (eds.) "Infections Caused by Anaerobic Bacteria," Jawetz, Melnick, & Adelberg's Medical Microbiology, 19th ed., pp. 257-262, Appleton & Lange, San Mateo, CA (1991).] As these organisms are relatively resistant to most commonly used antimicrobials, when a patient is treated with antibiotics, the other members of the normal gastrointestinal flora are suppressed and C. difficile flourishes, producing cytopathic toxins and enterotoxins. It has been found in 25% of cases of moderate diarrhea resulting from treatment with antibiotics, especially the cephalosporins, clindamycin, and ampicillin. [M.N. Swartz at 644.]

Importantly, C. difficile is commonly associated with nosocomial infections.

The organism is often present in the hospital and nursing home environments and may

be carried on the hands and clothing of hospital personnel who care for debilitated and immunocompromised patients. As many of these patients are being treated with antimicrobials or other chemotherapeutic agents, such transmission of *C. difficile* represents a significant risk factor for disease. (Engelkirk *et al.*, pp. 64-67.)

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C. difficile is associated with a range of diarrhetic illness, ranging from diarrhea alone to marked diarrhea and necrosis of the gastrointestinal mucosa with the accumulation of inflammatory cells and fibrin, which forms a pseudomembrane in the affected area. (Brooks et al.) It has been found in over 95% of pseudomembranous enterocolitis cases. (Swartz, at p. 644.) This occasionally fatal disease is characterized by diarrhea, multiple small colonic plaques, and toxic megacolon. (Swartz, at p. 644.) Although stool cultures are sometimes used for diagnosis, diagnosis is best made by detection of the heat labile toxins present in fecal filtrates from patients with enterocolitis due to C. difficile. (Swartz, at p. 644-645; and Brooks et al., at p. 260.) C. difficile toxins are cytotoxic for tissue/cell cultures and cause enterocolitis when injected intracecally into hamsters. (Swartz, at p. 644.)

The enterotoxicity of *C. difficile* is primarily due to the action of two toxins, designated A and B, each of approximately 300,000 in molecular weight. Both are potent cytotoxins, with toxin A possessing direct enterocytotoxic activity. [Lyerly et al., Infect. Immun. 60:4633 (1992).] Unlike toxin A of *C. perfringens*, an organism rarely associated with antimicrobial-associated diarrhea, the toxin of *C. difficile* is not a spore coat constituent and is not produced during sporulation. (Swartz, at p. 644.)

C. difficile toxin A causes hemorrhage, fluid accumulation and mucosal damage in rabbit ileal loops and appears to increase the uptake of toxin B by the intestinal mucosa. Toxin B does not cause intestinal fluid accumulation, but it is 1000 times more toxic than toxin A to tissue culture cells and causes membrane damage.

Although both toxins induce similar cellular effects such as actin disaggregation, differences in cell specificity occurs.

Both toxins are important in disease. [Borriello et al., Rev. Infect. Dis., 12(suppl. 2):S185 (1990); Lyerly et al., Infect. Immun., 47:349 (1985); and Rolfe, Infect. Immun., 59:1223 (1990).] Toxin A is thought to act first by binding to brush border receptors, destroying the outer mucosal layer, then allowing toxin B to gain access to the underlying tissue. These steps in pathogenesis would indicate that the production of neutralizing antibodies against toxin A may be sufficient in the prophylactic therapy of CDAD. However, antibodies against toxin B may be a necessary additional component for an effective therapeutic against later stage colonic disease. Indeed, it has been reported that animals require antibodies to both toxin A and toxin B to be completely protected against the disease. [Kim and Rolfe, Abstr. Ann. Meet. Am. Soc. Microbiol., 69:62 (1987).]

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C. difficile has also been reported to produce other toxins such as an enterotoxin different from toxins A and B [Banno et al., Rev. Infect. Dis., 6(Suppl. 1:S11-S20 (1984)], a low molecular weight toxin [Rihn et al., Biochem. Biophys. Res. Comm., 124:690-695 (1984)], a motility altering factor [Justus et al., Gastroenterol., 83:836-843 (1982)], and perhaps other toxins. Regardless, C. difficile gastrointestinal disease is of primary concern.

It is significant that due to its resistance to most commonly used antimicrobials, C. difficile is associated with antimicrobial therapy with virtually all antimicrobial agents (although most commonly ampicillin, clindamycin and cephalosporins). It is also associated with disease in patients undergoing chemotherapy with such compounds as methotrexate, 5-fluorouracil, cyclophosphamide, and doxorubicin. [S.M. Finegold et al., Clinical Guide to Anaerobic Infections, pp. 88-89, Star Publishing Co., Belmont, CA (1992).]

Treatment of *C. difficile* disease is problematic, given the high resistance of the organism. Oral metronidazole, bacitracin and vancomycin have been reported to be effective. (Finegold *et al.*, p. 89.) However there are problems associated with

treatment utilizing these compounds. Vancomycin is very expensive, some patients are unable to take oral medication, and the relapse rate is high (20-25%), although it may not occur for several weeks. *Id.*

C. difficile disease would be prevented or treated by neutralizing the effects of these toxins in the gastrointestinal tract. Thus, what is needed is an effective therapy against C. difficile toxin that is free of dangerous side effects, is available in large supply at a reasonable price, and can be safely delivered so that prophylactic application to patients at risk of developing pseudomembranous enterocolitis can be effectively treated.

10 DESCRIPTION OF THE DRAWINGS

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Figure 1 shows the reactivity of anti-C. botulinum IgY by Western blot.

Figure 2 shows the IgY antibody titer to C. botulinum type A toxoid in eggs, measured by ELISA.

Figure 3 shows the results of C. difficile toxin A neutralization assays.

Figure 4 shows the results of C. difficile toxin B neutralization assays.

Figure 5 shows the results of C. difficile toxin B neutralization assays.

Figure 6 is a restriction map of C. difficile toxin A gene, showing sequences of primers 1-4 (SEQ ID NOS:1-4).

Figure 7 is a Western blot of C. difficile toxin A reactive protein.

Figure 8 shows C. difficile toxin A expression constructs.

Figure 9 shows C. difficile toxin A expression constructs.

Figure 10 shows the purification of recombinant C. difficile toxin A.

Figure 11 shows the results of C. difficile toxin A neutralization assays with antibodies reactive to recombinant toxin A.

Figure 12 shows the results for a C. difficile toxin A neutralization plate.

Figure 13 shows the results for a C. difficile toxin A neutralization plate.

Figure 14 shows the results of recombinant C. difficile toxin A neutralization assays. Figure 15 shows C. difficile toxin A expression constructs. Figure 16 shows a chromatograph plotting absorbance at 280 nm against retention time for a pMA1870-680 IgY PEG preparation. Figure 17 shows two recombinant C. difficile toxin B expression constructs. Figure 18 shows C. difficile toxin B expression constructs. Figure 19 shows C. difficile toxin B expression constructs. Figure 20 shows C. difficile toxin B expression constructs. Figure 21 is an SDS-PAGE gel showing the purification of recombinant C. difficile toxin B fusion protein. Figure 22 is an SDS-PAGE gel showing the purification of two histidine-tagged recombinant C. difficile toxin B proteins. Figure 23 shows C. difficile toxin B expression constructs. Figure 24 is a Western blot of C. difficile toxin B reactive protein. Figure 25 shows C. botulinum type A toxin expression constructs; constructs used to provide C. botulinum or C. difficile sequences are also shown. Figure 26 is an SDS-PAGE gel stained with Coomaisse blue showing the purification of recombinant C. botulinum type A toxin fusion proteins. Figure 27 shows C. botulinum type A toxin expression constructs; constructs used to provide C. botulinum sequences are also shown. Figure 28 is an SDS-PAGE gel stained with Coomaisse blue showing the purification of pHisBot protein using the Ni-NTA resin. Figure 29 is an SDS-PAGE gel stained with Coomaisse blue showing the expression of pHisBot protein in BL21(DE3) and BL21(DE3)pLysS host cells. Figure 30 is an SDS-PAGE gel stained with Coomaisse blue showing the

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purification of pHisBot protein using a batch absorption procedure.

Figure 31 is an SDS-PAGE gel stained with Coomaisse blue showing the purification of pHisBot and pHisBot(native) proteins using a Ni-NTA column.

Figure 32 is an SDS-PAGE gel stained with Coomaisse blue showing the purification of pHisBotA protein expressed in pHisBotA(syn) kan lacIq T7/pACYCGro/BL21(DE3) cells using an IDA column.

Figure 33 is an SDS-PAGE gel stained with Coomaisse blue showing the purification of pHisBotA, pHisBotB and pHisBotE proteins by IDA chromatography followed by chromatography on S-100 to remove folding chaperones.

Figure 34 is an SDS-PAGE gel stained with Coomaisse blue showing the extracts derived from pHisBotB amp T7lac/BL21(DE3) cells before and after purification on a Ni-NTA column.

Figure 35 is an SDS-PAGE gel run under native conditions and stained with Coomaisse blue showing the removal of folding chaperones from IDA-purified BotB protein using a S-100 column.

Figure 36 is an SDS-PAGE gel stained with Coomaisse blue showing proteins that eluted during an imidazole step gradient applied to a IDA column containing a lysate of pHisBotB kan lacIq T7/pACYCGro/BL21(DE3) cells.

Figure 37 is an SDS-PAGE gel run under native conditions and stained with Coomaisse blue showing IDA-purified BotB protein before and after ultrafiltration.

Figure 38 is an SDS-PAGE gel stained with Coomaisse blue showing the purification of BotE protein using a NiNTA column.

Figure 39 is an SDS-PAGE gel stained with Coomaisse blue showing extracts derived from pHisBotA kan T7 lac/BL21(DE3) pLysS cells grown in fermentation culture.

Figure 40 is a chromatogram showing proteins present after IDA-purified BotE protein was applied to a S-100 column.

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DEFINITIONS

To facilitate understanding of the invention, a number of terms are defined below.

As used herein, the term "neutralizing" is used in reference to antitoxins, particularly antitoxins comprising antibodies, which have the ability to prevent the pathological actions of the toxin against which the antitoxin is directed.

As used herein, the term "overproducing" is used in reference to the production of clostridial toxin polypeptides in a host cell and indicates that the host cell is producing more of the clostridial toxin by virtue of the introduction of nucleic acid sequences encoding said clostridial toxin polypeptide than would be expressed by said host cell absent the introduction of said nucleic acid sequences. To allow ease of purification of toxin polypeptides produced in a host cell it is preferred that the host cell express or overproduce said toxin polypeptide at a level greater than 1 mg/liter of host cell culture.

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"A host cell capable of expressing a recombinant protein at a level greater than or equal to 5% of the total cellular protein" is a host cell in which the recombinant protein represents at least 5% of the total cellular protein. To determine what percentage of total cellular protein the recombinant protein represents, the following steps are taken. A total of 10 OD₆₀₀ units of recombinant host cells (e.g., 200 μl of cells at OD₆₀₀=50/ml) are removed (at a timepoint known to represent the peak of expression of the desired recombinant protein) to a 1.5 ml microfuge tube and pelleted for 2 min at maximum rpm in a microfuge. The pellets are resuspended in 1 ml of 50 mM NaHPO₄, 0.5 M NaCl, 40 mM imidazole buffer (pH 6.8) containing 1 mg/ml lysozyme. The samples are incubated for 20 min at room temperature and stored ON at -70°C. Samples are thawed completely at room temperature and sonicated 2 X 10 seconds with a Branson Sonifier 450 microtip probe at # 3 power setting. The samples are centrifuged for 5 min. at maximum rpm in a microfuge. An aliquot (20

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μl) of the protein sample is removed to 20 μl 2X sample buffer (this represents the total protein extract). The samples are heated to 95°C for 5 min, then cooled and 5 or 10 μl are loaded onto 12.5% SDS-PAGE gels. High molecular weight protein markers are also loaded to allow for estimation of the MW of identified recombinant proteins. After electrophoresis, protein is detected generally by staining with Coomassie blue and the stained gel is scanned using a densitometer to determine the percentage of protein present in each band. In this manner, the percentage of protein present in the band corresponding to the recombinant protein of interest may be determined. It is not necessary that Coomassie blue be employed for the detection of protein, a number of fluorescent dyes [e.g., Sypro orange S-6651 (Molecular Probes, Eugene, OR] may be employed and the stained gel scanned using a fluoroimager [e.g., Fluor Imager SI (Molecular Dynamics, Sunnyvale, CA)].

"A host cell capable of expressing a recombinant protein as a soluble protein at a level greater than or equal to 0.25% of the total soluble cellular protein" is a host cell in which the amount of soluble recombinant protein present represents at least 0.25% of the total cellular protein. As used herein "total soluble cellular protein" refers to a clarified PEI lysate prepared as described in Example 31(c)(iv). Briefly, cells are harvested following induction of expression of recombinant protein (at a point of maximal expression). The cells are resuspended in cell resuspension buffer (CRB: 50 mM NaPO₄, 0.5 M NaCl, 40 mM imidazole, pH 6.8) to create a 20% cell suspension (wet weight of cells/volume of CRB) and cell lysates are prepared as described in Example 31(c)(iv) (i.e., sonication or homogenization followed by centrifugation). The cell lysate is then flocculated utilizing polyethyleneimine (PEI) prior to centrifugation. PEI (a 2% solution in dH₂O, pH 7.5 with HCl) is added to the cell lysate to a final concentration of 0.2%, and stirred for 20 min at room temperature prior to centrifugation [8,500 rpm in JA10 rotor (Beckman) for 30 minutes at 4°C]. This treatment removes RNA, DNA and cell wall components, resulting in a clarified,

low viscosity lysate ("PEI clarified lysate"). The recombinant protein present in the PEI clarified lysate is then purified (e.g., by chromatography on an IDA column for his-tagged proteins). The amount of purified recombinant protein (i.e., the eluted protein) is divided by the concentration of protein present in the PEI clarified lysate (typically 8 mg/ml when using a 20% cell suspension as the starting material) and multiplied by 100 to determine what percentage of total soluble cellular protein is comprised of the soluble recombinant protein (see Example 33b).

As used herein, the term "fusion protein" refers to a chimeric protein containing the protein of interest (i.e., C. botulinum toxin A, B, C, D, E, F, or G and fragments thereof) joined to an exogenous protein fragment (the fusion partner which consists of a non-toxin protein). The fusion partner may enhance solubility of the C. botulinum protein as expressed in a host cell, may provide an affinity tag to allow purification of the recombinant fusion protein from the host cell or culture supernatant, or both. If desired, the fusion protein may be removed from the protein of interest (i.e., toxin protein or fragments thereof) prior to immunization by a variety of enzymatic or chemical means known to the art.

As used herein the term "non-toxin protein" or "non-toxin protein sequence" refers to that portion of a fusion protein which comprises a protein or protein sequence which is not derived from a bacterial toxin protein.

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The term "protein of interest" as used herein refers to the protein whose expression is desired within the fusion protein. In a fusion protein the protein of interest will be joined or fused with another protein or protein domain, the fusion partner, to allow for enhanced stability of the protein of interest and/or ease of purification of the fusion protein.

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As used herein, the term "maltose binding protein" refers to the maltose binding protein of *E. coli*. A portion of the maltose binding protein may be added to a protein of interest to generate a fusion protein; a portion of the maltose binding protein may

merely enhance the solubility of the resulting fusion protein when expressed in a bacterial host. On the other hand, a portion of the maltose binding protein may allow affinity purification of the fusion protein on an amylose resin.

As used herein, the term "poly-histidine tract" when used in reference to a fusion protein refers to the presence of two to ten histidine residues at either the amino- or carboxy-terminus of a protein of interest. A poly-histidine tract of six to ten residues is preferred. The poly-histidine tract is also defined functionally as being a number of consecutive histidine residues added to the protein of interest which allows the affinity purification of the resulting fusion protein on a nickel-chelate or IDA column.

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As used herein, the term "purified" or "to purify" refers to the removal of contaminants from a sample. For example, antitoxins are purified by removal of contaminating non-immunoglobulin proteins; they are also purified by the removal of immunoglobulin that does not bind toxin. The removal of non-immunoglobulin proteins and/or the removal of immunoglobulins that do not bind toxin results in an increase in the percent of toxin-reactive immunoglobulins in the sample. In another example, recombinant toxin polypeptides are expressed in bacterial host cells and the toxin polypeptides are purified by the removal of host cell proteins; the percent of recombinant toxin polypeptides is thereby increased in the sample. Additionally, the recombinant toxin polypeptides are purified by the removal of host cell components such as lipopolysaccharide (e.g., endotoxin).

The term "recombinant DNA molecule" as used herein refers to a DNA molecule which is comprised of segments of DNA joined together by means of molecular biological techniques.

The term "recombinant protein" or "recombinant polypeptide" as used herein refers to a protein molecule which is expressed from a recombinant DNA molecule.

The term "native protein" as used herein refers to a protein which is isolated from a natural source as opposed to the production of a protein by recombinant means.

As used herein the term "portion" when in reference to a protein (as in "a portion of a given protein") refers to fragments of that protein. The fragments may range in size from four amino acid residues to the entire amino acid sequence minus one amino acid.

As used herein "soluble" when in reference to a protein produced by recombinant DNA technology in a host cell is a protein which exists in solution in the cytoplasm of the host cell; if the protein contains a signal sequence the soluble protein is exported to the periplasmic space in bacteria hosts and is secreted into the culture medium in eucaryotic cells capable of secretion or by bacterial host possessing the appropriate genes (i.e., the kil gene). In contrast, an insoluble protein is one which exists in denatured form inside cytoplasmic granules (called inclusion bodies) in the host cell. High level expression (i.e., greater than 10-20 mg recombinant protein/liter of bacterial culture) of recombinant proteins often results in the expressed protein being found in inclusion bodies in the bacterial host cells. A soluble protein is a protein which is not found in an inclusion body inside the host cell or is found both in the cytoplasm and in inclusion bodies and in this case the protein may be present at high or low levels in the cytoplasm.

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A distinction is drawn between a soluble protein (i.e., a protein which when expressed in a host cell is produced in a soluble form) and a "solubilized" protein. An insoluble recombinant protein found inside an inclusion body may be solubilized (i.e., rendered into a soluble form) by treating purified inclusion bodies with denaturants such as guanidine hydrochloride, urea or sodium dodecyl sulfate (SDS). These denaturants must then be removed from the solubilized protein preparation to allow the recovered protein to renature (refold). Not all proteins will refold into an active conformation after solubilization in a denaturant and removal of the denaturant. Many

proteins precipitate upon removal of the denaturant. SDS may be used to solubilize inclusion bodies and will maintain the proteins in solution at low concentration. However, dialysis will not always remove all of the SDS (SDS can form micelles which do not dialyze out); therefore, SDS-solubilized inclusion body protein is soluble but not refolded.

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A distinction is drawn between proteins which are soluble (i.e., dissolved) in a solution devoid of significant amounts of ionic detergents (e.g., SDS) or denaturants (e.g., urea, guanidine hydrochloride) and proteins which exist as a suspension of insoluble protein molecules dispersed within the solution. A soluble protein will not be removed from a solution containing the protein by centrifugation using conditions sufficient to remove bacteria present in a liquid medium (i.e., centrifugation at 12,000 x g for 4-5 minutes). For example, to test whether two proteins, protein A and protein B, are soluble in solution, the two proteins are placed into a solution selected from the group consisting of PBS-NaCl (PBS containing 0.5 M NaCl), PBS-NaCl containing 0.2% Tween 20, PBS, PBS containing 0.2% Tween 20, PBS-C (PBS containing 2 mM CaCl₃), PBS-C containing either 0.1 or 0.5 % Tween 20, PBS-C containing either 0.1 or 0.5% NP-40, PBS-C containing either 0.1 or 0.5% Triton X-100, PBS-C containing 0.1% sodium deoxycholate. The mixture containing proteins A and B is then centrifuged at 5000 x g for 5 minutes. The supernatant and pellet formed by centrifugation are then assayed for the presence of protein A and B. If protein A is found in the supernatant and not in the pellet [except for minor amounts (i.e., less than 10%) as a result of trapping, protein is said to be soluble in the solution tested. If the majority of protein B is found in the pellet (i.e., greater than 90%), then protein B is said to exist as a suspension in the solution tested.

As used herein, the term "therapeutic amount" refers to that amount of antitoxin required to neutralize the pathologic effects of one or more clostridial toxins in a subject.

The term "pyrogen" as used herein refers to a fever-producing substance. Pyrogens may be endogenous to the host (e.g., prostaglandins) or may be exogenous compounds (e.g., bacterial endo- and exotoxins, nonbacterial compounds such as antigens and certain steroid compounds, etc.). The presence of pyrogen in a pharmaceutical solution may be detected using the U.S. Pharmacopeia (USP) rabbit fever test (United States Pharmacopeia, Vol. XXII (1990) United States Pharmacopeial Convention, Rockville, MD, p. 151).

The term "endotoxin" as used herein refers to the high molecular weight complexes associated with the outer membrane of gram-negative bacteria. Unpurified endotoxin contains lipids, proteins and carbohydrates. Highly purified endotoxin does not contain protein and is referred to as lipopolysaccharide (LPS). Because unpurified endotoxin is of concern in the production of pharmaceutical compounds (e.g., proteins produced in E. coli using recombinant DNA technology), the term endotoxin as used herein refers to unpurified endotoxin. Bacterial endotoxin is a well known pyrogen.

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As used herein, the term "endotoxin-free" when used in reference to a composition to be administered parenterally (with the exception of intrathecal administration) to a host means that the dose to be delivered contains less than 5 EU/kg body weight [FDA Guidelines for Parenteral Drugs (December 1987)]. Assuming a weight of 70 kg for an adult human, the dose must contain less than 350 EU to meet FDA Guidelines for parenteral administration. Endotoxin levels are measured herein using the Limulus Amebocyte Lysate (LAL) test (Limulus Amebocyte Lysate PyrochromeTM, Associates of Cape Cod, Inc. Woods Hole, MA). To measure endotoxin levels in preparations of recombinant proteins, 0.5 ml of a solution comprising 0.5 mg of purified recombinant protein in 50 mM NaPO₄, pH 7.0, 0.3M NaCl and 10% glycerol is used in the LAL assay according to the manufacturer's instructions for the endpoint chromogenic without diazo-coupling method [the specific components of the buffer containing recombinant protein to be analyzed in the LAL

test are not important; any buffer having a neutral pH may be employed (see for example, alternative buffers employed in Examples 34, 40 and 45)]. Compositions containing less than or equal to than 250 endotoxin units (EU)/mg of purified recombinant protein are herein defined as "substantially endotoxin-free." Preferably the composition contains less than or equal to 100, and most preferably less than or equal to 60, (EU)/mg of purified recombinant protein. Typically, administration of bacterial toxins or toxoids to adult humans for the purpose of vaccination involves doses of about 10-500 µg protein/dose. Therefore, administration of 10-500 µg of a purified recombinant protein to a 70 kg human, wherein said purified recombinant protein preparation contains 60 EU/mg protein, results in the introduction of only 0.6 to 30 EU (i.e., 0.2 to 8.6% of the maximum allowable endotoxin burden per parenteral dose). Administration of 10-500 µg of a purified recombinant protein to a 70 kg human, wherein said purified recombinant protein preparation contains 250 EU/mg protein, results in the introduction of only 2.5 to 125 EU (i.e., 0.7 to 36% of the maximum allowable endotoxin burden per parenteral dose).

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The LAL test is accepted by the U.S. FDA as a means of detecting bacterial endotoxins (21 C.F.R. §§ 660.100 -105). Studies have shown that the LAL test is equivalent or superior to the USP rabbit pyrogen test for the detection of endotoxin and thus the LAL test can be used as a surrogate for pyrogenicity studies in animals [F.C. Perason, *Pyrogens: endotoxins, LAL testing and depyrogenation*, Marcel Dekker, New York (1985), pp.150-155]. The FDA Bureau of Biologics accepts the LAL assay in place of the USP rabbit pyrogen test so long as the LAL assay utilized is shown to be as sensitive as, or more sensitive as the rabbit test [Fed. Reg., 38, 26130 (1980)].

The term "monovalent" when used in reference to a clostridial vaccine refers to a vaccine which is capable of provoking an immune response in a host animal directed against a single type of clostridial toxin. For example, if immunization of a host with C. botulinum type A toxin vaccine induces antibodies in the immunized host which

protect against a challenge with type A toxin but not against challenge with type B, C, D, E, F or G toxins, then the type A vaccine is said to be monovalent. In contrast, a "multivalent" vaccine provokes an immune response in a host animal directed against several (i.e., more than one) clostridial toxins. For example, if immunization of a host with a vaccine comprising C. botulinum type A and B toxins induces the production of antibodies which protect the host against a challenge with both type A and B toxin, the vaccine is said to be multivalent (in particular, this hypothetical vaccine is bivalent).

As used herein the term "immunogenically-effective amount" refers to that amount of an immunogen required to invoke the production of protective levels of antibodies in a host upon vaccination.

The term "protective level", when used in reference to the level of antibodies induced upon immunization of the host with an immunogen which comprises a bacterial toxin, means a level of circulating antibodies sufficient to protect the host from challenge with a lethal dose of the toxin.

As used herein the terms "protein" and "polypeptide" refer to compounds comprising amino acids joined via peptide bonds and are used interchangeably.

The terms "toxin" and "neurotoxin" when used in reference to toxins produced by members (i.e., species and strains) of the genus Clostridium are used interchangeably and refer to the proteins which are poisonous to nerve tissue.

The term "receptor-binding domain" when used in reference to a *C. botulinum* toxin refers to the carboxy-terminal portion of the heavy chain (H_C or the C fragment) of the toxin which is presumed to be responsible for the binding of the active toxin (i.e., the derivative toxin comprising the H and L chains joined via disulfide bonds) to receptors on the surface of synaptosomes. The receptor-binding domain for *C. botulinum* type A toxin is defined herein as comprising amino acid residues 861 through 1296 of SEQ ID NO:28. The receptor-binding domain for *C. botulinum* type B toxin is defined herein as comprising amino acid residues 848 through 1291 of SEQ

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ID NO:40 (strain Eklund 17B). The receptor-binding domain of C. botulinum type C1 toxin is defined herein as comprising amino acid residues 856 through 1291 of SEQ ID NO:60. The receptor-binding domain of C. botulinum type D toxin is defined herein as comprising amino acid residues 852 through 1276 of SEQ ID NO:66. The receptor-binding domain of C. botulinum type E toxin is defined herein as comprising amino acid residues 835 through 1250 of SEQ ID NO:50 (Beluga strain). The receptor-binding domain of C. botulinum type F toxin is defined herein as comprising amino acid residues 853 through 1274 of SEQ ID NO:71. The receptor-binding domain of C. botulinum type G toxin is defined herein as comprising amino acid residues 853 through 1297 of SEQ ID NO:77. Within a given serotype, small variations in the primary amino acid sequence of the botulinal toxins isolated from different strains has been reported [Whelan et al. (1992), supra and Minton (1995) Curr. Top. Microbiol. Immunol. 195:161-194]. The present invention contemplates fusion proteins comprising the receptor-binding domain of C. botulinum toxins from serotypes A-G including the variants found among different strains within a given serotype. The receptor-binding domains listed above are used as the prototype for each strain within a serotype. Fusion proteins containing an analogous region from a strain other than the prototype strain are encompassed by the present invention.

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Fusion proteins comprising the receptor binding domain (i.e., C fragment) of botulinal toxins may include amino acid residues located beyond the termini of the domains defined above. For example, the pHisBotB protein contains amino acid residues 846-1291 of SEQ ID NO:40; this fusion protein thus comprises the receptor-binding domain for *C. botulinum* type B toxin as defined above (i.e., Ile-848 through Glu-1291). Similarly, pHisBotE contains amino acid residues 827-1252 of SEQ ID NO:50 and pHisBotG contains amino acid residues 851-1297 of SEQ ID NO:77. Thus, both pHisBotE and pHisBotG fusion proteins contain a few amino acids located beyond the N-terminus of the defined receptor-binding domain.

The terms "native gene" or "native gene sequences" are used to indicate DNA sequences encoding a particular gene which contain the same DNA sequences as found in the gene as isolated from nature. In contrast, "synthetic gene sequences" are DNA sequences which are used to replace the naturally occurring DNA sequences when the naturally occurring sequences cause expression problems in a given host cell. For example, naturally-occurring DNA sequences encoding codons which are rarely used in a host cell may be replaced (e.g., by site-directed mutagenesis) such that the synthetic DNA sequence represents a more frequently used codon. The native DNA sequence and the synthetic DNA sequence will preferably encode the same amino acid sequence.

SUMMARY OF THE INVENTION

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The present invention relates to the production of polypeptides derived from toxins particularly in recombinant host cells. In one embodiment, the present invention provides a host cell containing a recombinant expression vector, said vector encoding a protein comprising at least a portion of a Clostridium botulinum toxin, said toxin selected from the group consisting of type B toxin and type E toxin. The present invention is not limited by the nature of sequences encoding portions of the C. botulinum toxin. These sequences may be derived from the native gene sequences or alternatively they may comprise synthetic gene sequences. Synthetic gene sequences are employed when expression of the native gene sequences is problematic in a given host cell (e.g., when the native gene sequences contain sequences resembling yeast transcription termination signals and the desired host cell is a yeast cell).

In one embodiment, the host cell is capable of expressing the recombinant C. botulinum toxin protein at a level greater than or equal to 2% to 40% of the total cellular protein and preferably at a level greater than or equal to 5% of the total cellular protein. In another embodiment, the host cell is capable of expressing the recombinant C. botulinum toxin protein as a soluble protein at a level greater than or

equal to 0.25% of the total cellular protein and preferably at a level greater than or equal to 0.25% to 10% of the total cellular protein.

The present invention is not limited by the nature of the host cell employed for the production of recombinant *C. botulinum* toxin proteins. In a preferred embodiment, the host cell is an *E. coli* cell. In another preferred embodiment, the host cell is an insect cell; particularly preferred insect host cells are *Spodoptera frugiperda* (Sf9) cells. In another preferred embodiment, the host cell is a yeast cell; particularly preferred yeast cells are *Pichia pastoris* cells.

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In another embodiment, the invention provides a host cell containing a recombinant expression vector, said vector encoding a fusion protein comprising a non-toxin protein sequence and at least a portion of a Clostridium botulinum toxin, said toxin selected from the group consisting of type B toxin and type E toxin. The invention is not limited by the nature of the portion of the Clostridium botulinum toxin selected. In a preferred embodiment, the portion of the toxin comprises the receptor binding domain (i.e., the C fragment of the toxin). The present invention is not limited by the nature of the non-toxin protein sequence employed. In a preferred embodiment, the non-toxin protein sequence comprises a poly-histidine tract. A number of alternative fusion tags or fusion partners are known to the art (e.g., MBP, GST, protein A, etc.) and may be employed for the production of fusion proteins comprising a portion of a botulinal toxin.

The present invention further provides a vaccine comprising a fusion protein, said fusion protein comprising a non-toxin protein sequence and at least a portion of a Clostridium botulinum toxin, said toxin selected from the group consisting of type B toxin and type E toxin. The vaccine may be a monovalent vaccine (i.e., containing only a toxin B fusion protein or a toxin E fusion protein), a bivalent vaccine (i.e., containing both a toxin B fusion protein and a toxin E fusion protein) or a trivalent or higher valency vaccine. In a preferred embodiment, the toxin B fusion protein and/or

toxin E fusion protein is combined with a fusion protein comprising a non-toxin protein sequence and at least a portion of Clostridium botulinum type A toxin. The present invention is not limited by the nature of the portion of the Clostridium botulinum toxin selected. In a preferred embodiment, the portion of the toxin comprises the receptor binding domain (i.e., the C fragment of the toxin). The present invention is not limited by the nature of the non-toxin protein sequence employed. In a preferred embodiment, the non-toxin protein sequence comprises a poly-histidine tract. A number of alternative fusion tags or fusion partners are known to the art (e.g., MBP, GST, protein A, etc.) and may be employed for the generation of fusion proteins comprising vaccines. When a fusion partner (i.e., the non-toxin protein sequence) is employed for the production of a recombinant C. botulinal toxin protein, the fusion partner may be removed from the recombinant C. botulinal toxin protein if desired (i.e., prior to administration of the protein to a subject) using a variety of methods known to the art (e.g., digestion of fusion proteins containing FactorXa or thrombin recognition sites with the appropriate enzyme). A number of the pETHis vectors employed herein provide an N-terminal his-tag followed by a FactorXa cleavage site (see Example 28a); the botulinal C fragment sequences follow the FactorXa site and thus, FactorXa can be used to remove the his-tag from the botulinal fusion protein. In a preferred embodiment, the vaccine is substantially endotoxin-free.

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The present invention is not limited by the method employed for the generation of vaccine comprising fusion proteins comprising a non-toxin protein sequence and at least a portion of a Clostridium botulinum toxin. The fusion proteins may be produced by recombinant DNA means using either native or synthetic gene sequences expressed in a host cell. The present invention is not limited to the production of vaccines using recombinant host cells; cell free in vitro transcription/translation systems may be employed for the expression of the nucleic acid constructs encoding the fusion proteins of the present invention. An example of such a cell-free system is the commercially

available TnTTM Coupled Reticulocyte Lysate System (Promega Corporation, Madison, WI). Alternatively, the fusion proteins of the present invention may be generated by synthetic means (i.e., peptide synthesis).

The present invention further provides a method of generating antibody directed against a Clostridium botulinum toxin comprising: a) providing in any order: i) an antigen comprising a fusion protein comprising a non-toxin protein sequence and at least a portion of a Clostridium botulinum toxin, said toxin selected from the group consisting of type B toxin and type E toxin, and ii) a host; and b) immunizing the host with the antigen so as to generate an antibody. In a preferred embodiment, the antigen used to immunize the host also contains a fusion protein comprising a non-toxin protein sequence and at least a portion of Clostridium botulinum type A toxin. The present invention is not limited by the nature of the portion of the Clostridium botulinum toxin selected. In a preferred embodiment, the portion of the toxin comprises the receptor binding domain (i.e., the C fragment of the toxin). The present invention is not limited by the nature of the non-toxin protein sequence employed. In a preferred embodiment, the non-toxin protein sequence comprises a poly-histidine tract. A number of alternative fusion tags or fusion partners are known to the art (e.g., MBP, GST, protein A, etc.) and may be employed for the generation of fusion proteins comprising vaccines. When a fusion partner (i.e., the non-toxin protein sequence) is employed for the production of a recombinant C. botulinal toxin protein, the fusion partner may be removed from the recombinant C. botulinal toxin protein if desired (i.e., prior to administration of the protein to a subject) using a variety of methods known to the art (e.g., digestion of fusion proteins containing FactorXa or thrombin recognition sites with the appropriate enzyme).

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The present invention is not limited by the nature of the host employed for the production of the antibodies of the invention. In a preferred embodiment, the host is a mammal, preferably a human. The antibodies of the present invention may be

generated using non-mammalian hosts such as birds, preferably chickens. In a preferred embodiment the method of the present invention further comprised the step c) of collecting the antibodies from the host. In yet another embodiment, the method of the present invention further comprises the step d) of purifying the antibodies.

The present invention further provides antibodies raised according to the above methods.

The present invention further contemplates multivalent vaccines comprising at least two recombinant C. botulinum toxin proteins derived from the group consisting of C. botulinum serotypes A, B, C, D, E, F, and G. The invention contemplates bivalent, trivalent, quadravalent, pentavalent, heptavalent and septivalent vaccines comprising recombinant C. botulinum toxin proteins. Preferably the recombinant C. botulinum toxin proteins toxin protein comprises the receptor binding domain (i.e., C fragment) of the toxin.

DESCRIPTION OF THE INVENTION

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The present invention contemplates vaccinating humans and other animals with polypeptides derived from *C. botulinum* neurotoxins which are substantially endotoxin-free. These botulinal peptides are also useful for the production of antitoxin. Antibotulinal toxin antitoxin is useful for the treatment of patients effected by or at risk of symptoms due to the action of *C. botulinum* toxins. The organisms, toxins and individual steps of the present invention are described separately below.

I. Clostridium Species, Clostridial Diseases And Associated Toxins

A preferred embodiment of the method of the present invention is directed toward obtaining antibodies against *Clostridium* species, their toxins, enzymes or other metabolic by-products, cell wall components, or synthetic or recombinant versions of any of these compounds. It is contemplated that these antibodies will be produced by immunization of humans or other animals. It is not intended that the present invention

be limited to any particular toxin or any species of organism. In one embodiment, toxins from all Clostridium species are contemplated as immunogens. Examples of these toxins include the neuraminidase toxin of C. butyricum, C. sordellii toxins HT and LT, toxins A, B, C, D, E, F, and G of C. botulinum and the numerous C. perfringens toxins. In one preferred embodiment, toxins A, B, and E of C. botulinum are contemplated as immunogens. Table 2 above lists various Clostridium species, their toxins and some antigens associated with disease.

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TABLE 2
Clostridial Toxins

10	Organism	Toxins and Disease-Associated Antigens
	C. botulinum	A, B, C ₁ , C ₂ , D, E, F, G
15	C. butyricum	Neuraminidase
	C. difficile	A, B, Enterotoxin (not A nor B), Motility Altering Factor, Low Molecular Weight Toxin, Others
	C. perfringens	α, β, ε, ι, γ, δ, ν, θ, κ, λ, μ, υ
	C. sordelli/ C. bifermentans	ΗΤ, LΤ, α, β, γ
	C. novyi	α, β, γ, δ, ε, ζ, ν, θ
	C. septicum	α, β, γ, δ
	C. histolyticum	α, β, γ, δ, ε plus additional enzymes
20	C. chawoei	α, β, γ, δ

It is not intended that antibodies produced against one toxin will only be used against that toxin. It is contemplated that antibodies directed against one toxin (e.g., C. perfringens type A enterotoxin) may be used as an effective therapeutic against one or more toxin(s) produced by other members of the genus Clostridium or other toxin producing organisms (e.g., Bacillus cereus, Staphylococcus aureus, Streptococcus mutans, Acinetobacter calcoaceticus, Pseudomonas aeruginosa, other Pseudomonas

species, etc.). It is further contemplated that antibodies directed against the portion of the toxin which binds to mammalian membranes (e.g., C. perfringens enterotoxin A) can also be used against other organisms. It is contemplated that these membrane binding domains are produced synthetically and used as immunogens.

II. Obtaining Antibodies In Non-Mammals

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A preferred embodiment of the method of the present invention for obtaining antibodies involves immunization. However, it is also contemplated that antibodies could be obtained from non-mammals without immunization. In the case where no immunization is contemplated, the present invention may use non-mammals with preexisting antibodies to toxins as well as non-mammals that have antibodies to whole organisms by virtue of reactions with the administered antigen. An example of the latter involves immunization with synthetic peptides or recombinant proteins sharing epitopes with whole organism components.

In a preferred embodiment, the method of the present invention contemplates immunizing non-mammals with bacterial toxin(s). It is not intended that the present invention be limited to any particular toxin. In one embodiment, toxin from all clostridial bacteria sources (see Table 2) are contemplated as immunogens. Examples of these toxins are C. butyricum neuraminidase toxin, toxins A, B, C, D, E, F, and G from C. botulinum, C. perfringens toxins α , β , ε , and ε , and ε , and ε and ε . In a preferred embodiment, C. botulinum toxins A, B, C, D, E, and F (or fragments thereof) are contemplated as immunogens.

A particularly preferred embodiment involves the use of bacterial toxin protein or fragments of toxin proteins produced by molecular biological means (i.e., recombinant toxin proteins). In a preferred embodiment, the immunogen comprises the receptor-binding domain (i.e., the ~50 kD carboxy-terminal portion of the heavy chain; also referred to as the C fragment) of C. botulinum serotype A neurotoxin produced by recombinant DNA technology. In another preferred embodiment, the immunogen comprises the receptor-binding domain of C. botulinum serotype B

neurotoxin produced by recombinant DNA technology. In yet another preferred embodiment, the immunogen comprises the receptor-binding domain region of C. botulinum serotype E neurotoxin produced by recombinant DNA technology. In yet another preferred embodiment, the immunogen comprises the receptor-binding domain region of C. botulinum serotype C1 neurotoxin produced by recombinant DNA technology. In yet another preferred embodiment, the immunogen comprises the receptor-binding domain region of C. botulinum serotype C2 neurotoxin produced by recombinant DNA technology. In yet another preferred embodiment, the immunogen comprises the receptor-binding domain region of C. botulinum serotype D neurotoxin produced by recombinant DNA technology. In yet another preferred embodiment, the immunogen comprises the receptor-binding domain region of C. botulinum serotype F neurotoxin produced by recombinant DNA technology. In yet another preferred embodiment, the immunogen comprises the receptor-binding domain region of C. botulinum serotype G neurotoxin produced by recombinant DNA technology. In a preferred embodiment, the recombinant botulinal toxin proteins are expressed as fusion proteins (e.g., as histidine-tagged proteins). In a still further preferred embodiment, the immunogen is a multivalent vaccine comprising the receptor-binding domain region of C. botulinum toxin from two or more toxins selected from the group consisting of type A, type B, type C (including C1 and C2), type D, type E, and type F toxin.

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When immunization is used, the preferred non-mammal is from the class Aves. All birds are contemplated (e.g., duck, ostrich, emu, turkey, etc.). A preferred bird is a chicken. Importantly, chicken antibody does not fix mammalian complement. [See H.N. Benson et al., J. Immunol. 87:616 (1961).] Thus, chicken antibody will normally not cause a complement-dependent reaction. [A.A. Benedict and K. Yamaga, "Immunoglobulins and Antibody Production in Avian Species," in Comparative Immunology (J.J. Marchaloni, ed.), pp. 335-375, Blackwell, Oxford (1966).] Thus, the preferred antitoxins of the present invention will not exhibit complement-related side effects observed with antitoxins known presently.

When birds are used, it is contemplated that the antibody will be obtained from either the bird serum or the egg. A preferred embodiment involves collection of the antibody from the egg. Laying hens transport immunoglobulin to the egg yolk ("IgY") in concentrations equal to or exceeding that found in serum. [See R. Patterson et al., J. Immunol. 89:272 (1962); and S.B. Carroll and B.D. Stollar, J. Biol. Chem. 258:24 (1983).] In addition, the large volume of egg yolk produced vastly exceeds the volume of serum that can be safely obtained from the bird over any given time period. Finally, the antibody from eggs is purer and more homogeneous; there is far less non-immunoglobulin protein (as compared to serum) and only one class of immunoglobulin is transported to the yolk.

When considering immunization with toxins, one may consider modification of the toxins to reduce the toxicity. In this regard, it is not intended that the present invention be limited by immunization with modified toxin. Unmodified ("native") toxin is also contemplated as an immunogen.

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It is also not intended that the present invention be limited by the type of modification -- if modification is used. The present invention contemplates all types of toxin modification, including chemical and heat treatment of the toxin. The preferred modification, however, is formaldehyde treatment.

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It is not intended that the present invention be limited to a particular mode of immunization; the present invention contemplates all modes of immunization, including subcutaneous, intramuscular, intraperitoneal, and intravenous or intravascular injection, as well as *per os* administration of immunogen.

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The present invention further contemplates immunization with or without adjuvant. (Adjuvant is defined as a substance known to increase the immune response to other antigens when administered with other antigens.) If adjuvant is used, it is not intended that the present invention be limited to any particular type of adjuvant -- or that the same adjuvant, once used, be used all the time. While the present invention contemplates all types of adjuvant, whether used separately or in combinations, the preferred use of adjuvant is the use of Complete Freund's Adjuvant followed sometime later with Incomplete Freund's Adjuvant. Another preferred use of adjuvant is the use

of Gerbu Adjuvant. The invention also contemplates the use of RIBI fowl adjuvant and Quil A adjuvant.

When immunization is used, the present invention contemplates a wide variety of immunization schedules. In one embodiment, a chicken is administered toxin(s) on day zero and subsequently receives toxin(s) in intervals thereafter. It is not intended that the present invention be limited by the particular intervals or doses. Similarly, it is not intended that the present invention be limited to any particular schedule for collecting antibody. The preferred collection time is sometime after day 100.

Where birds are used and collection of antibody is performed by collecting eggs, the eggs may be stored prior to processing for antibody. It is preferred that eggs be stored at 4°C for less than one year.

It is contemplated that chicken antibody produced in this manner can be bufferextracted and used analytically. While unpurified, this preparation can serve as a reference for activity of the antibody prior to further manipulations (e.g., immunoaffinity purification).

III. Increasing The Effectiveness Of Antibodies

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When purification is used, the present invention contemplates purifying to increase the effectiveness of both non-mammalian antitoxins and mammalian antitoxins. Specifically, the present invention contemplates increasing the percent of toxin-reactive immunoglobulin. The preferred purification approach for avian antibody is polyethylene glycol (PEG) separation.

The present invention contemplates that avian antibody be initially purified using simple, inexpensive procedures. In one embodiment, chicken antibody from eggs is purified by extraction and precipitation with PEG. PEG purification exploits the differential solubility of lipids (which are abundant in egg yolks) and yolk proteins in high concentrations of PEG 8000. [Polson et al., Immunol. Comm. 9:495 (1980).] The technique is rapid, simple, and relatively inexpensive and yields an immunoglobulin fraction that is significantly purer in terms of contaminating non-immunoglobulin proteins than the comparable ammonium sulfate fractions of

mammalian sera and horse antibodies. The majority of the PEG is removed from the precipitated chicken immunoglobulin by treatment with ethanol. Indeed, PEG-purified antibody is sufficiently pure that the present invention contemplates the use of PEG-purified antitoxins in the passive immunization of intoxicated humans and animals.

IV. Treatment

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The present invention contemplates antitoxin therapy for humans and other animals intoxicated by bacterial toxins. A preferred method of treatment is by intravenous administration of anti-boutlinal antitoxin; oral administration is also contemplated for other clostridial antitoxins.

A. Dosage Of Antitoxin

It was noted by way of background that a balance must be struck when administering currently available antitoxin which is usually produced in large animals such as horses; sufficient antitoxin must be administered to neutralize the toxin, but not so much antitoxin as to increase the risk of untoward side effects. These side effects are caused by: i) patient sensitivity to foreign (e.g, horse) proteins; ii) anaphylactic or immunogenic properties of non-immunoglobulin proteins; iii) the complement fixing properties of mammalian antibodies; and/or iv) the overall burden of foreign protein administered. It is extremely difficult to strike this balance when, as noted above, the degree of intoxication (and hence the level of antitoxin therapy needed) can only be approximated.

The present invention contemplates significantly reducing side effects so that this balance is more easily achieved. Treatment according to the present invention contemplates reducing side effects by using PEG-purified antitoxin from birds.

In one embodiment, the treatment of the present invention contemplates the use of PEG-purified antitoxin from birds. The use of yolk-derived, PEG-purified antibody as antitoxin allows for the administration of: 1) non(mammalian)-complement-fixing, avian antibody; 2) a less heterogeneous mixture of non-immunoglobulin proteins; and 3) less total protein to deliver the equivalent weight of active antibody present in

currently available antitoxins. The non-mammalian source of the antitoxin makes it useful for treating patients who are sensitive to horse or other mammalian sera.

B. Delivery Of Antitoxin

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Although it is not intended to limit the route of delivery, the present invention contemplates a method for antitoxin treatment of bacterial intoxication in which delivery of antitoxin is oral. In one embodiment, antitoxin is delivered in a solid form (e.g., tablets). In an alternative embodiment antitoxin is delivered in an aqueous solution. When an aqueous solution is used, the solution has sufficient ionic strength to solubilize antibody protein, yet is made palatable for oral administration. The delivery solution may also be buffered (e.g., carbonate buffer pH 9.5) which can neutralize stomach acids and stabilize the antibodies when the antibodies are administered orally. In one embodiment the delivery solution is an aqueous solution. In another embodiment the delivery solution is a nutritional formula. Preferably, the delivery solution is infant formula. Yet another embodiment contemplates the delivery of lyophilized antibody encapsulated or microencapsulated inside acid-resistant compounds.

Methods of applying enteric coatings to pharmaceutical compounds are well known to the art [companies specializing in the coating of pharmaceutical compounds are available; for example, The Coating Place (Verona, WI) and AAI (Wilmington, NC)]. Enteric coatings which are resistant to gastric fluid and whose release (i.e., dissolution of the coating to release the pharmaceutical compound) is pH dependent are commercially available [for example, the polymethacrylates Eudragit® L and Eudragit® S (Röhm GmbH)]. Eudragit® S is soluble in intestinal fluid from pH 7.0; this coating can be used to microencapsulate lyophilized antitoxin antibodies and the particles are suspended in a solution having a pH above or below pH 7.0 for oral administration. The microparticles will remain intact and undissolved until they reached the intestines where the intestinal pH would cause them to dissolve thereby releasing the antitoxin.

The invention contemplates a method of treatment which can be administered for treatment of acute intoxication. In one embodiment, antitoxin is administered orally in either a delivery solution or in tablet form, in therapeutic dosage, to a subject intoxicated by the bacterial toxin which served as immunogen for the antitoxin.

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The invention also contemplates a method of treatment which can be administered prophylactically. In one embodiment, antitoxin is administered orally, in a delivery solution, in therapeutic dosage, to a subject, to prevent intoxication of the subject by the bacterial toxin which served as immunogen for the production of antitoxin. In another embodiment, antitoxin is administered orally in solid form such as tablets or as microencapsulated particles. Microencapsulation of lyophilized antibody using compounds such as Eudragit® (Rohm GmbH) or polyethylene glycol, which dissolve at a wide range of pH units, allows the oral administration of solid antitoxin in a liquid form (i.e., a suspension) to recipients unable to tolerate administration of tablets (e.g., children or patients on feeding tubes). In one preferred embodiment the subject is a child. In another embodiment, antibody raised against whole bacterial organism is administered orally to a subject, in a delivery solution, in therapeutic dosage.

V. Vaccines Against Clostridial Species

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The invention contemplates the generation of mono- and multivalent vaccines for the protection of an animal (particularly humans) against several clostridial species. Of particular interest are vaccines which stimulate the production of a humoral immune response to *C. botulinum, C. tetani and C. difficile* in humans. The antigens comprising the vaccine preparation may be native or recombinantly produced toxin proteins from the clostridial species listed above. When toxin proteins are used as immunogens they are generally modified to reduce the toxicity. This modification may be by chemical or genetic (*i.e.*, recombinant DNA technology) means. In general genetic detoxification (*i.e.*, the expression of nontoxic fragments in a host cell) is preferred as the expression of nontoxic fragments in a host cell precludes the presence

of intact, active toxin in the final preparation. However, when chemical modification is desired, the preferred toxin modification is formaldehyde treatment.

The invention contemplates that recombinant *C. botulinum* toxin proteins be used as antigens in mono- and multivalent vaccine preparations. Soluble, substantially endotoxin-free recombinant *C. botulinum* toxin proteins derived from serotypes A, B and E may be used individually (*i.e.*, as mono-valent vaccines) or in combination (*i.e.*, as a multi-valent vaccine). In addition, the recombinant *C. botulinum* toxin proteins derived from serotpes A, B and E may be used in conjunction with either recombinant or native toxins or toxoids from other serotypes of *C. botulinum*, *C. difficile* and *C. tetani* as antigens for the preparation of these mono- and multivalent vaccines. It is contemplated that, due to the structural similarity of *C. botulinum* and *C. tetani* toxin proteins, a vaccine comprising *C. difficile* and *botulinum* toxin proteins (native or recombinant or a mixture thereof) be used to stimulate an immune response against *C. botulinum*, *C. tetani and C. difficile*.

The present invention further contemplates multi-valent vaccines comprising two or more botulinal toxin proteins selected from the group comprising recombinant C. botulinum toxin proteins derived from serotypes A, B, C (including C1 and C2), D, E, F and G.

The adverse consequences of exposure to botulinal toxin would be avoided by immunization of subjects at risk of exposure to the toxin with nontoxic preparations which confer immunity such as chemically or genetically detoxified toxin.

Vaccines which confer immunity against one or more of the toxin types A, B, E, F and G would be useful as a means of protecting humans from the deleterious effects of those C. botulinum toxins known to affect man. Indeed as the possibility exists that humans could be exposed to any of the seven serotypes of C. botulinum toxin (e.g., during biological warfare or the production of toxin in a laboratory setting), multivalent vaccines capable of conferring immunity against toxin types A-G (including both C1 and C2 toxins) would be useful for the protection of humans.

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Vaccines which confer immunity against one or more of the toxin types C, D and E would be useful for veterinary applications.

The botulinal neurotoxin is synthesized as a single polypeptide chain which is processed into a heavy (H; ~100 kD) and a light (L; ~50 kD) chain by cleavage with proteolytic enzymes; these two chains are held together via disulfide bonds in the active toxin (referred to as derivative toxin) [B.R. DasGupta and H. Sugiyama, Biochem. Biophys. Res. Commun. 48:108 (1972); reviewed in B.R. DasGupta, J. Physiol. 84:220 (1990), H. Sugiyama, Microbiol. Rev. 44:419 (1980) and C.L. Hatheway, Clin. Microbiol. Rev. 3:66 (1990)]. The heavy chain of the active toxin is cleaved by trypsin to produce two fragments termed H_c (also referred to as H₁ or C) and H_N (also referred to as H₂ or B). The H_C fragment (~46 kD) comprises the carboxy end of the H chain. The H_N fragment (~49 kD) comprises the animo end and remains attached to the L chain (H_NL) . Neither H_C or H_NL is toxic. H_C competes with whole derivative toxin for binding to synaptosomes and therefore H_C is said to contain the receptor binding site. The H_C and H_N fragments of botulinal toxin are analogous to the fragments C and B of tetanus toxin which are produced by papain cleavage. The C fragment of tetanus toxin has been shown to be responsible for the binding of tetanus toxin to purified gangliosides and neuronal cells [Halpern and Loftus, J. Biol. Chem. 288:11188 (1993)].

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Antisera raised against purified preparations of isolated botulinal H and L chains have been shown to protect mice against the lethal effects of the toxin; however, the effectiveness of the two antisera differ with the anti-H sera being more potent (H. Sugiyama, supra). While the different botulinal toxins show structural similarity to one another, the different serotypes are reported to be immunologically distinct (i.e., sera raised against one toxin type does not cross-react to a significant degree with other types). Thus, the generation of multivalent vaccines may require the use of more than one type of toxin.

C. botulinum toxin genes from all seven serotypes have been cloned and sequenced (Minton (1995), supra); in addition, partial amino acid sequence is available

for a number of *C. botulinum* toxins isolated from different strains within a given serotype. The *C. botulinum* toxins contain about 1250-1300 amino acid residues. On the DNA level, the overall degree of homology between *C. botulinum* serotypes A, B, C, D and E toxins averages between 50 and 60% identity with a greater degree of homology being found between H chain-encoding regions than between those encoding L chains [Whelan *et al.* (1992) Appl. Environ. Microbiol. 58:2345]. The degree of identity between *C. botulinum* toxins on the amino acid level reflects the level of DNA sequence homology. The most divergent area of DNA and amino acid sequence is found within the carboxy-terminal area of the various *C. botulinum* H chain genes. This portion of the toxin (*i.e.*, H_C or the C fragment) plays a major role in cell binding. As toxin from different serotypes is thought to bind to distinct cell receptor molecules, it is not surprising that the toxins diverge significantly over this region.

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Within a given serotype, small variations in the primary amino acid sequence of the botulinal toxins isolated from different strains has been reported [Whelan et al. (1992), supra and Minton (1995), supra]. The present invention contemplates fusion proteins comprising portions of C. botulinum toxins from serotypes A-G including the variants found among different strains within a given serotype. The present invention provides oligonucleotide primers which may be used to amplify the C fragment or receptor-binding region of the toxin gene from various strains of C. botulinum serotype A, serotype B, serotype C (C1 and C2), serotype D, serotype E, serotype F and serotype G. A large number of different strains of C. botulinum serotype A, serotype B, serotype C, serotype D serotype E and serotype F are available from the American Type Culture Collection (ATCC; Rockville, MD). For example, the ATCC provides the following: Type A strains: 174 (ATCC 3502), 457 (ATCC 17862), and NCTC 7272 (ATCC 19397); Type B strains: 34 (ATCC 439), 62A (ATCC 7948), NCA 213 B (ATCC 7949), 13114 (ATCC 8083), 3137 (ATCC 17780), 1347 (ATCC 17841), 2017 (ATCC 17843), 2217 (ATCC 17844), 2254 (ATCC 17845) and VP 1731 (ATCC 25765); Type C strains: 2220 (ATCC 17782), 2239 (ATCC 17783), 2223 (ATCC 17784; a type C-β strain; C-β strains produce C2 toxin), 662 (ATCC 17849; a type C-

α strain; C-α strains produce mainly C1 toxin and a small amount of C2 toxin), 2021 (ATCC 17850; a type C-α strain) and VPI 3803 (ATCC 25766); Type D strains: ATCC 9633, 2023 (ATCC 17851), and VPI 5995 (ATCC 27517); Type E strains: ATCC 43181, 36208 (ATCC 9564), 2231 (ATCC 17786), 2229 (ATCC 17852), 2279 (ATCC 17854) and 2285 (ATCC 17855) and Type F strains: 202F (ATCC 23387), VPI 4404 (ATCC 25764), VPI 2382 (ATCC 27321) and Langeland (ATCC 35415). Type G strain, 113/30 (NCFB 3012) may be obtained from the National Collection of Food Bacteria (NCFB, AFRC Institute of Food Research, Reading, United Kingdom).

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Purification methods have been reported for native toxin types A, B, C, D, E, and F [reviewed in G. Sakaguchi, Pharmac. Ther. 19:165 (1983)]. As the different botulinal toxins are structurally related, the invention contemplates the expression of any of the botulinal toxins (e.g., types A-G) as soluble recombinant fusion proteins.

In particular, methods for purification of the type A botulinum neurotoxin have been developed [L.J. Moberg and H. Sugiyama, Appl. Environ. Microbiol. 35:878 (1978)]. Immunization of hens with detoxified purified protein results in the generation of neutralizing antibodies [B.S. Thalley et al., in Botulinum and Tetanus Neurotoxins, B.R. DasGupta, ed., Plenum Press, New York (1993), p. 467].

The currently available *C. botulinum* pentavalent vaccine comprising chemically inactivated (*i.e.*, formaldehyde treated) type A, B, C, D and E toxins is not adequate. The efficacy is variable (in particular, only 78% of recipients produce protective levels of anti-type B antibodies following administration of the primary series) and immunization is painful (deep subcutaneous inoculation is required for administration), with adverse reactions being common (moderate to severe local reactions occur in approximately 6% of recipients upon initial injection; this number rises to approximately 11% of individuals who receive booster injections) [Informational Brochure for the Pentavalent (ABCDE) Botulinum Toxoid, Centers for Disease Control]. Preparation of this vaccine is dangerous as active toxin must be handled by laboratory workers.

In general, chemical detoxification of bacterial toxins using agents such as formaldehyde, glutaraldehyde or hydrogen peroxide is not optimal for the generation of vaccines or antitoxins. A delicate balance must be struck between too much and too little chemical modification. If the treatment is insufficient, the vaccine may retain residual toxicity. If the treatment is too excessive, the vaccine may lose potency due to destruction of native immunogenic determinants. Another major limitation of using botulinal toxoids for the generation of antitoxins or vaccines is the high production expense. For the above reasons, the development of methods for the production of nontoxic but immunogenic *C. botulinum* toxin proteins is desirable.

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The C. botulinum and C. tetanus toxin proteins have similar structures [reviewed in E.J. Schantz and E.A. Johnson, Microbiol. Rev. 56:80 (1992)]. The carboxy-terminal 50 kD fragment of the tetanus toxin heavy chain (fragment C) is released by papain cleavage and has been shown to be non-toxic and immunogenic. Recombinant tetanus toxin fragment C has been developed as a candidate vaccine antigen [A.J. Makoff et al., Bio/Technology 7:1043 (1989)]. Mice immunized with recombinant tetanus toxin fragment C were protected from challenge with lethal doses of tetanus toxin. No studies have demonstrated that the recombinant tetanus fragment C protein confers immunity against other botulinal toxins such as the C. botulinum toxins.

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Recombinant tetanus fragment C has been expressed in E. coli (A.J. Makoff et al., Bio/Technology, supra and Nucleic Acids Res. 17:10191 (1989); J.L. Halpern et al., Infect. Immun. 58:1004 (1990)], yeast [M.A. Romanos et al., Nucleic Acids Res. 19:1461 (1991)] and baculovirus [I.G. Charles et al., Infect. Immun. 59:1627 (1991)]. Synthetic tetanus toxin genes had to be constructed to facilitate expression in yeast (M.A. Romanos et al., supra) and E. coli [A.J. Makoff et al., Nucleic Acids Res., supra], due to the high A-T content of the tetanus toxin gene sequences. High A-T content is a common feature of clostridial genes (M.R. Popoff et al., Infect. Immun. 59:3673 (1991); H.F. LaPenotiere et al., in Botulinum and Tetanus Neurotoxins, B.R. DasGupta, ed., Plenum Press, New York (1993), p. 463] which creates expression

difficulties in *E. coli* and yeast due primarily to altered codon usage frequency and fortuitous polyadenylation sites, respectively.

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The C fragment of the C. botulinum type A neurotoxin heavy chain has been evaluated as a vaccine candidate. The C. botulinum type A neurotoxin gene has been cloned and sequenced [D.E. Thompson et al., Eur. J. Biochem. 189:73 (1990)]. The C fragment of the type A toxin was expressed as either a fusion protein comprising the botulinal C fragment fused with the maltose binding protein (MBP) or as a native protein [H.F. LaPenotiere et al., (1993) supra, H.F. LaPenotiere et al., Toxicon. 33:1383 (1995) and Middlebrook and Brown (1995), Curr. Top. Microbiol. Immunol. 195:89-122]. The plasmid construct encoding the native protein was reported to be unstable, while the fusion protein was expressed primarily in inclusion bodies as insoluble protein. Immunization of mice with crudely purified MBP fusion protein resulted in protection against IP challenge with 3 LD50 doses of toxin [LaPenotiere et al., (1993) and (1995), supra]. However, this recombinant C. botulinum type A toxin C fragment/MBP fusion protein is not a suitable immunogen for the production of vaccines as it is expressed as an insoluble protein in E. coli. Furthermore, this recombinant C. botulinum type A toxin C fragment/MBP fusion protein was not shown to be substantially free of endotoxin contamination. Experience with recombinant C. botulinum type A toxin C fragment/MBP fusion proteins shows that the presence of the MBP on the fusion protein greatly complicates the removal of endotoxin from preparations of the recombinant fusion protein (see Ex. 24, infra). Expression of a synthetic gene encoding C. botulinum type A toxin C fragment as a soluble protein excreted from insect cells has been reported [Middlebrook and Brown (1995), supra]; no details regarding the level of expression achieved or the presence of endotoxin or other pyrogens were provided. Like the insoluble protein expressed in E. coli, immunization with the recombinant protein produced in insect cells was reported to protect mice from challenge with C. botulinum toxin A.

Inclusion body protein must be solubilized prior to purification and/or administration to a host. The harsh treatment of inclusion body protein needed to

accomplish this solubilization may reduce the immunogenicity of the purified protein. Ideally, recombinant proteins to be used as vaccines are expressed as soluble proteins at high levels (i.e., greater than or equal to about 0.75% of total cellular protein) in E. coli or other host cells (e.g., yeast, insect cells, etc.). This facilitates the production and isolation of sufficient quantities of the immunogen in a highly purified form (i.e., substantially free of endotoxin or other pyrogen contamination). The ability to express recombinant toxin proteins as soluble proteins in E. coli is advantageous due to the low cost of growth compared to insect or mammalian tissue culture cells.

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The C. botulinum type B neurotoxin gene has been cloned and sequenced from two strains of C. botulinum type B [Whelan et al. (1992) Appl. Environ. Microbiol. 58:2345 (Danish strain) and Hutson et al. (1994) Curr. Microbiol. 28:101 (Eklund 17B strain)]. The nucleotide sequence of the toxin gene derived from the Eklund 17B strain (ATCC 25765) is available from the EMBL/GenBank sequence data banks under the accession number X71343; the nucleotide sequence of the coding region is listed in SEQ ID NO:39. The amino acid sequence of the C. botulinum type B neurotoxin derived from the strain Eklund 17B is listed in SEQ ID NO:40. The nucleotide sequence of the C. botulinum serotype B toxin gene derived from the Danish strain is listed in SEQ ID NO:41. The amino acid sequence of the C. botulinum type B neurotoxin derived from the Danish strain is listed in SEQ ID NO:42.

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The C. botulinum type B neurotoxin gene is synthesized as a single polypeptide chain which is processed to form a dimer composed of a light and a heavy chain linked via disulfide bonds. The light chain is responsible for pharmacological activity (i.e., inhibition of the release of acetylcholine at the neuromuscular junction). The N-terminal portion of the heavy chain is thought to mediate channel formation while the C-terminal portion mediates toxin binding; the type B neurotoxin has been reported to exist as a mixture of predominantly single chain with some double chain (Whelan et al., supra). The 50 kD carboxy-terminal portion of the heavy chain is referred to as the C fragment or the H_C domain. The present invention reports for the first time, the

expression of the C fragment of C. botulinum type B toxin in heterologous hosts (e.g., E. coli).

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The C. botulinum type E neurotoxin gene has been cloned and sequenced from a number of different strains [Poulet et al. (1992) Biochem. Biophys. Res. Commun. 183:107; Whelan et al. (1992) Eur. J. Biochem. 204:657; and Fujii et al. (1993) J. Gen. Microbiol. 139:79]. The nucleotide sequence of the type E toxin gene is available from the EMBL sequence data bank under accession numbers X62089 (strain Beluga) and X62683 (strain NCTC 11219); the nucleotide sequence of the coding region (strain Beluga) is listed in SEQ ID NO:45. The amino acid sequence of the C. botulinum type E neurotoxin derived from strain Beluga is listed in SEQ ID NO:46. The type E neurotoxin gene is synthesized as a single polypeptide chain which may be converted to a double-chain form (i.e., a heavy chain and a light chain) by cleavage with trypsin; unlike the type A neurotoxin, the type E neurotoxin exists essentially only in the single-chain form. The 50 kD carboxy-terminal portion of the heavy chain is referred to as the C fragment or the H_C domain. The present invention reports for the first time, the expression of the C fragment of C. botulinum type E toxin in heterologous hosts (e.g., E. coli).

The C. botulinum type C1, D, F and G neurotoxin genes have been cloned and sequenced. The nucleotide and amino acid sequences of these genes and toxins are provided herein. The invention provides methods for the expression of the C fragment from each of these toxin genes in heterologous hosts and the purification of the resulting recombinant proteins.

The subject invention provides methods which allow the production of soluble C. botulinum toxin proteins in economical host cells (e.g., E. coli). In addition the subject invention provides methods which allow the production of soluble botulinal toxin proteins in yeast and insect cells. Further, methods for the isolation of purified soluble C. botulinum toxin proteins which are suitable for immunization of humans and other animals are provided. These soluble, purified preparations of C. botulinum

toxin proteins provide the basis for improved vaccine preparations and facilitate the production of antitoxin.

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When recombinant clostridial toxin proteins produced in gram-negative bacteria (e.g., E. coli) are used as vaccines, they are purified to remove endotoxin prior to administration to a host animal. In order to vaccinate a host, an immunogenically-effective amount of purified substantially endotoxin-free recombinant clostridial toxin protein is administered in any of a number of physiologically acceptable carriers known to the art. When administered for the purpose of vaccination, the purified substantially endotoxin-free recombinant clostridial toxin protein may be used alone or in conjunction with known adjutants, including potassium alum, aluminum phosphate, aluminum hydroxide, Gerbu adjuvant (GmDP; C.C. Biotech Corp.), RIBI adjuvant (MPL; RIBI Immunochemical Research, Inc.), QS21 (Cambridge Biotech). The alum and aluminum-based adjutants are particularly preferred when vaccines are to be administered to humans; however, any adjuvant approved for use in humans may be employed. The route of immunization may be nasal, oral, intramuscular, intraperitoneal or subcutaneous.

The invention contemplates the use of soluble, substantially endotoxin-free preparations of fusion proteins comprising the C fragment of the C. botulinum type A, B, C, D, E, F, and G toxin as vaccines. In one embodiment, the vaccine comprises the C fragment of either the C. botulinum type A, B, C, D, E, F, or G toxin and a polyhistidine tract (also called a histidine tag). In a particularly preferred embodiment, a fusion protein comprising the histidine tagged C fragment is expressed using the pET series of expression vectors (Novagen). The pET expression system utilizes a vector containing the T7 promoter which encodes the fusion protein and a host cell which can be induced to express the T7 DNA polymerase (i.e., a DE3 host strain). The production of C fragment fusion proteins containing a histidine tract is not limited to the use of a particular expression vector and host strain. Several commercially available expression vectors and host strains can be used to express the C fragment protein sequences as a fusion protein containing a histidine tract (For example, the pQE series (pQE-8, 12, 16, 17, 18, 30, 31, 32, 40, 41, 42, 50, 51, 52, 60 and 70) of

expression vectors (Qiagen) which are used with the host strains M15[pREP4] (Qiagen) and SG13009[pREP4] (Qiagen) can be used to express fusion proteins containing six histidine residues at the amino-terminus of the fusion protein). Furthermore a number of commercially available expression vectors which provide a histidine tract also provide a protease cleavage site between the histidine tract and the protein of interest (e.g., botulinal toxin sequences). Cleavage of the resulting fusion protein with the appropriate protease will remove the histidine tag from the protein of interest (e.g., botulinal toxin sequences) (see Example 28a, infra). Removal of the histidine tag may be desirable prior to administration of the recombinant botulinal toxin protein to a subject (e.g., a human).

VI. Detection Of Toxin

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The invention contemplates detecting bacterial toxin in a sample. The term "sample" in the present specification and claims is used in its broadest sense. On the one hand it is meant to include a specimen or culture. On the other hand, it is meant to include both biological and environmental samples.

Biological samples may be animal, including human, fluid, solid (e.g., stool) or tissue; liquid and solid food products and ingredients such as dairy items, vegetables, meat and meat by-products, and waste. Environmental samples include environmental material such as surface matter, soil, water and industrial samples, as well as samples obtained from food and dairy processing instruments, apparatus, equipment, disposable and non-disposable items. These examples are not to be construed as limiting the sample types applicable to the present invention.

The invention contemplates detecting bacterial toxin by a competitive immunoassay method that utilizes recombinant toxin A and toxin B proteins, antibodies raised against recombinant bacterial toxin proteins. A fixed amount of the recombinant toxin proteins are immobilized to a solid support (e.g., a microtiter plate) followed by the addition of a biological sample suspected of containing a bacterial toxin. The biological sample is first mixed with affinity-purified or PEG fractionated

antibodies directed against the recombinant toxin protein. A reporter reagent is then added which is capable of detecting the presence of antibody bound to the immobilized toxin protein. The reporter substance may comprise an antibody with binding specificity for the antitoxin attached to a molecule which is used to identify the presence of the reporter substance. If toxin is present in the sample, this toxin will compete with the immobilized recombinant toxin protein for binding to the anti-recombinant antibody thereby reducing the signal obtained following the addition of the reporter reagent. A control is employed where the antibody is not mixed with the sample. This gives the highest (or reference) signal.

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The invention also contemplates detecting bacterial toxin by a "sandwich" immunoassay method that utilizes antibodies directed against recombinant bacterial toxin proteins. Affinity-purified antibodies directed against recombinant bacterial toxin proteins are immobilized to a solid support (e.g., microtiter plates). Biological samples suspected of containing bacterial toxins are then added followed by a washing step to remove substantially all unbound antitoxin. The biological sample is next exposed to the reporter substance, which binds to antitoxin and is then washed free of substantially all unbound reporter substance. The reporter substance may comprise an antibody with binding specificity for the antitoxin attached to a molecule which is used to identify the presence of the reporter substance. Identification of the reporter substance in the biological tissue indicates the presence of the bacterial toxin.

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It is also contemplated that bacterial toxin be detected by pouring liquids (e.g., soups and other fluid foods and feeds including nutritional supplements for humans and other animals) over immobilized antibody which is directed against the bacterial toxin. It is contemplated that the immobilized antibody will be present in or on such supports as cartridges, columns, beads, or any other solid support medium. In one embodiment, following the exposure of the liquid to the immobilized antibody, unbound toxin is substantially removed by washing. The exposure of the liquid is then exposed to a reporter substance which detects the presence of bound toxin. In a preferred embodiment the reporter substance is an enzyme, fluorescent dye, or radioactive compound attached to an antibody which is directed against the toxin (i.e.,

in a "sandwich" immunoassay). It is also contemplated that the detection system will be developed as necessary (e.g., the addition of enzyme substrate in enzyme systems; observation using fluorescent light for fluorescent dye systems; and quantitation of radioactivity for radioactive systems).

EXPERIMENTAL

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The following examples serve to illustrate certain preferred embodiments and aspects of the present invention and are not to be construed as limiting the scope thereof.

In the disclosure which follows, the following abbreviations apply: °C (degrees Centigrade); rpm (revolutions per minute); BBS-Tween (borate buffered saline containing Tween); BSA (bovine serum albumin); ELISA (enzyme-linked immunosorbent assay); CFA (complete Freund's adjuvant); IFA (incomplete Freund's adjuvant); IgG (immunoglobulin G); IgY (immunoglobulin Y); IM (intramuscular); IP (intraperitoneal); IV (intravenous or intravascular); SC (subcutaneous); H₂O (water); HCl (hydrochloric acid); LD₁₀₀ (lethal dose for 100% of experimental animals); aa (amino acid); HPLC (high performance liquid chromatography); kD (kilodaltons); gm (grams); µg (micrograms); mg (milligrams); ng (nanograms); µl (microliters); ml (milliliters); mm (millimeters); nm (nanometers); µm (micrometer); M (molar); mM (millimolar); MW (molecular weight); sec (seconds); min(s) (minute/minutes); hr(s) (hour/hours); MgCl₂ (magnesium chloride); NaCl (sodium chloride); Na₂CO₃ (sodium carbonate); OD₂₈₀ (optical density at 280 nm); OD₆₀₀ (optical density at 600 nm); PAGE (polyacrylamide gel electrophoresis); PBS [phosphate buffered saline (150 mM NaCl, 10 mM sodium phosphate buffer, pH 7.2)]; PEG (polyethylene glycol); PMSF (phenylmethylsulfonyl fluoride); SDS (sodium dodecyl sulfate); Tris (tris(hydroxymethyl)aminomethane); Ensure® (Ensure®, Ross Laboratories, Columbus OH); Enfamil® (Enfamil®, Mead Johnson); w/v (weight to volume); v/v (volume to volume); Amicon (Amicon, Inc., Beverly, MA); Amresco (Amresco, Inc., Solon, OH); ATCC (American Type Culture Collection, Rockville, MD); BBL (Baltimore Biologics Laboratory, (a division of Becton Dickinson), Cockeysville, MD); Becton Dickinson

(Becton Dickinson Labware, Lincoln Park, NJ); BioRad (BioRad, Richmond, CA); Biotech (C-C Biotech Corp., Poway, CA); Charles River (Charles River Laboratories, Wilmington, MA); Cocalico (Cocalico Biologicals Inc., Reamstown, PA); CytRx (CytRx Corp., Norcross, GA); Falcon (e.g. Baxter Healthcare Corp., McGaw Park, IL and Becton Dickinson); FDA (Federal Food and Drug Administration); Fisher Biotech (Fisher Biotech, Springfield, NJ); GIBCO (Grand Island Biologic Company/BRL, Grand Island, NY); Gibco-BRL (Life Technologies, Inc., Gaithersburg, MD); Harlan Sprague Dawley (Harlan Sprague Dawley, Inc., Madison, WI); Mallinckrodt (a division of Baxter Healthcare Corp., McGaw Park, IL); Millipore (Millipore Corp., Marlborough, MA); New England Biolabs (New England Biolabs, Inc., Beverly, MA); Novagen (Novagen, Inc., Madison, WI); Pharmacia (Pharmacia, Inc., Piscataway, NJ); Qiagen (Qiagen, Chatsworth, CA); Sasco (Sasco, Omaha, NE); Showdex (Showa Denko America, Inc., New York, NY); Sigma (Sigma Chemical Co., St., Louis, MO); Sterogene (Sterogene, Inc., Arcadia, CA); Tech Lab (Tech Lab, Inc., Blacksburg, VA); and Vaxcell (Vaxcell, Inc., a subsidiary of CytRX Corp., Norcross, GA).

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When a recombinant protein is described in the specification it is referred to in a short-hand manner by the amino acids in the toxin sequence present in the recombinant protein rounded to the nearest 10. For example, the recombinant protein pMB1850-2360 contains amino acids 1852 through 2362 of the *C. difficile* toxin B protein. The specification gives detailed construction details for all recombinant proteins such that one skilled in the art will know precisely which amino acids are present in a given recombinant protein.

EXAMPLE 1

Production Of High-Titer Antibodies To Clostridium difficile Organisms In A Hen

Antibodies to certain pathogenic organisms have been shown to be effective in treating diseases caused by those organisms. It has not been shown whether antibodies can be raised, against *Clostridium difficile*, which would be effective in treating

infection by this organism. Accordingly, C. difficile was tested as immunogen for production of hen antibodies.

To determine the best course for raising high-titer egg antibodies against whole C. difficile organisms, different immunizing strains and different immunizing concentrations were examined. The example involved (a) preparation of the bacterial immunogen, (b) immunization, (c) purification of anti-bacterial chicken antibodies, and (d) detection of anti-bacterial antibodies in the purified IgY preparations.

a) Preparation Of Bacterial Immunogen

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C. difficile strains 43594 (serogroup A) and 43596 (serogroup C) were originally obtained from the ATCC. These two strains were selected because they represent two of the most commonly-occurring serogroups isolated from patients with antibiotic-associated pseudomembranous colitis. [Delmee et al., J. Clin. Microbiol., 28(10):2210 (1990).] Additionally, both of these strains have been previously characterized with respect to their virulence in the Syrian hamster model for C. difficile infection. [Delmee et al., J. Med Microbiol., 33:85 (1990).]

The bacterial strains were separately cultured on brain heart infusion agar for 48 hours at 37°C in a Gas Pack 100 Jar (BBL, Cockeysville, MD) equipped with a Gas Pack Plus anaerobic envelope (BBL). Forty-eight hour cultures were used because they produce better growth and the organisms have been found to be more cross-reactive with respect to their surface antigen presentation. The greater the degree of cross-reactivity of our IgY preparations, the better the probability of a broad range of activity against different strains/serogroups. [Toma et al., J. Clin. Microbiol., 26(3):426 (1988).]

The resulting organisms were removed from the agar surface using a sterile dacron-tip swab, and were suspended in a solution containing 0.4% formaldehyde in PBS, pH 7.2. This concentration of formaldehyde has been reported as producing good results for the purpose of preparing whole-organism immunogen suspensions for the generation of polyclonal anti-C. difficile antisera in rabbits. [Delmee et al., J. Clin.

Microbiol., 21:323 (1985); Davies et al., Microbial Path., 9:141 (1990).] In this manner, two separate bacterial suspensions were prepared, one for each strain. The two suspensions were then incubated at 4°C for 1 hour. Following this period of formalin-treatment, the suspensions were centrifuged at 4,200 x g for 20 min., and the resulting pellets were washed twice in normal saline. The washed pellets, which contained formalin-treated whole organisms, were resuspended in fresh normal saline such that the visual turbidity of each suspension corresponded to a #7 McFarland standard. [M.A.C. Edelstein, "Processing Clinical Specimens for Anaerobic Bacteria: Isolation and Identification Procedures," in S.M. Finegold et al (eds.)., Bailey and Scott's Diagnostic Microbiology, pp. 477-507, C.V. Mosby Co., (1990). The preparation of McFarland nephelometer standards and the corresponding approximate number of organisms for each tube are described in detail at pp. 172-173 of this volume.] Each of the two #7 suspensions was then split into two separate volumes. One volume of each suspension was volumetrically adjusted, by the addition of saline, to correspond to the visual turbidity of a #1 McFarland standard. [Id.] The #1 suspensions contained approximately 3 x 10⁸ organisms/ml, and the #7 suspensions contained approximately 2 x 109 organisms/ml. [Id.] The four resulting concentrationadjusted suspensions of formalin-treated C. difficile organisms were considered to be "bacterial immunogen suspensions." These suspensions were used immediately after preparation for the initial immunization. [See section (b).]

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The formalin-treatment procedure did not result in 100% non-viable bacteria in the immunogen suspensions. In order to increase the level of killing, the formalin concentration and length of treatment were both increased for subsequent immunogen preparations, as described below in Table 3. (Although viability was decreased with the stronger formalin treatment, 100% inviability of the bacterial immunogen suspensions was not reached.) Also, in subsequent immunogen preparations, the formalin solutions were prepared in normal saline instead of PBS. At day 49, the day of the fifth immunization, the excess volumes of the four previous bacterial

immunogen suspensions were stored frozen at -70°C for use during all subsequent immunizations.

b) Immunization

For the initial immunization, 1.0 ml volumes of each of the four bacterial immunogen suspensions described above were separately emulsified in 1.2 ml volumes of CFA (GIBCO). For each of the four emulsified immunogen suspensions, two fourmonth old White Leghorn hens (pre-laying) were immunized. (It is not necessary to use pre-laying hens; actively-laying hens can also be utilized.) Each hen received a total volume of approximately 1.0 ml of a single emulsified immunogen suspension via four injections (two subcutaneous and two intramuscular) of approximately 250 µl per site. In this manner, a total of four different immunization combinations, using two hens per combination, were initiated for the purpose of evaluating both the effect of immunizing concentration on egg yolk antibody (IgY) production, and interstrain cross-reactivity of IgY raised against heterologous strains. The four immunization groups are summarized in Table 3.

TABLE 3
Immunization Groups

minum zaton Groups			
Group Designation	Immunizing Strain	Approximate Immunizing Dose	
CD 43594, #1	C. difficile strain 43594	1.5 × 10 ⁴ organisms/hen	
CD 43594, #7	11 11	1.0 × 10° organisms/hen	
CD 43596, #1	C. difficile strain 43596	1.5 × 10 ^s organisms/hen	
CD 43596, #7	11 11	1.0 × 10° organisms/hen	

The time point for the first series of immunizations was designated as "day zero." All subsequent immunizations were performed as described above except that the bacterial immunogen suspensions were emulsified using IFA (GIBCO) instead of CFA, and for the later time point immunization, the stored frozen suspensions were

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used instead of freshly-prepared suspensions. The immunization schedule used is listed in Table 4.

TABLE 4
Immunization Schedule

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Day Of Immunization	Formalin-Treatment	Immunogen Preparation Used
0	1%, 1 hr.	freshly-prepared
14	1%, overnight	n u
21	1%, overnight	61 01
35	1%, 48 hrs.	19 11
49	1%, 72 hrs.	и и
70	11 11	stored frozen
85	11 11	17 10
105	11 11	11 14

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c) Purification Of Anti-Bacterial Chicken Antibodies

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Groups of four eggs were collected per immunization group between days 80 and 84 post-initial immunization, and chicken immunoglobulin (IgY) was extracted according to a modification of the procedure of A. Polson et al., Immunol. Comm., 9:495 (1980). A gentle stream of distilled water from a squirt bottle was used to separate the yolks from the whites, and the yolks were broken by dropping them through a funnel into a graduated cylinder. The four individual yolks were pooled for each group. The pooled, broken yolks were blended with 4 volumes of egg extraction buffer to improve antibody yield (egg extraction buffer is 0.01 M sodium phosphate, 0.1 M NaCl, pH 7.5, containing 0.005% thimerosal), and PEG 8000 (Amresco) was added to a concentration of 3.5%. When all the PEG dissolved, the protein precipitates that formed were pelleted by centrifugation at 13,000 × g for 10 minutes. The supernatants were decanted and filtered through cheesecloth to remove the lipid layer, and the PEG was added to the supernatants to a final concentration of 12% (the supernatants were assumed to contain 3.5% PEG). After a second centrifugation, the supernatants were discarded and the pellets were centrifuged a final time to extrude the

remaining PEG. These crude IgY pellets were then dissolved in the original yolk volume of egg extraction buffer and stored at 4°C. As an additional control, a preimmune IgY solution was prepared as described above, using eggs collected from unimmunized hens.

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d) Detection Of Anti-Bacterial Antibodies In The Purified IgY Preparations

In order to evaluate the relative levels of specific anti-C. difficile activity in the IgY preparations described above, a modified version of the whole-organism ELISA procedure of N.V. Padhye et al., J. Clin. Microbiol. 29:99-103 (1990) was used. Frozen organisms of both C. difficile strains described above were thawed and diluted to a concentration of approximately 1×10^7 organisms/ml using PBS, pH 7.2. In this way, two separate coating suspensions were prepared, one for each immunizing strain. Into the wells of 96-well microtiter plates (Falcon, Pro-Bind Assay Plates) were placed 100 µl volumes of the coating suspensions. In this manner, each plate well received a total of approximately 1×10^6 organisms of one strain or the other. The plates were then incubated at 4°C overnight. The next morning, the coating suspensions were decanted, and all wells were washed three times using PBS. In order to block nonspecific binding sites, 100 µl of 0.5% BSA (Sigma) in PBS was then added to each well, and the plates were incubated for 2 hours at room temperature. The blocking solution was decanted, and 100 µl volumes of the IgY preparations described above were initially diluted 1:500 with a solution of 0.1% BSA in PBS, and then serially diluted in 1:5 steps. The following dilutions were placed in the wells: 1:500, 1:2,500, 1:62,5000, 1:312,500, and 1:1,562,500. The plates were again incubated for 2 hours at room temperature. Following this incubation, the IgY-containing solutions were decanted, and the wells were washed three times using BBS-Tween (0.1 M boric acid, 0.025 M sodium borate, 1.0 M NaCl, 0.1% Tween-20), followed by two washes using PBS-Tween (0.1% Tween-20), and finally, two washes using PBS only. To each well, 100 µl of a 1:750 dilution of rabbit anti-chicken IgG (whole-molecule)-alkaline phosphatase conjugate (Sigma) (diluted in 0.1% BSA in PBS) was added. The plates

were again incubated for 2 hours at room temperature. The conjugate solutions were decanted and the plates were washed as described above, substituting 50 mM Na₂CO₃, pH 9.5 for the PBS in the final wash. The plates were developed by the addition of 100 µl of a solution containing 1 mg/ml para-nitrophenyl phosphate (Sigma) dissolved in 50 mM Na₂CO₃, 10 mM MgCl₂, pH 9.5 to each well, and incubating the plates at room temperature in the dark for 45 minutes. The absorbance of each well was measured at 410 nm using a Dynatech MR 700 plate reader. In this manner, each of the four IgY preparations described above was tested for reactivity against both of the immunizing *C. difficile* strains; strain-specific, as well as cross-reactive activity was determined.

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Table 5 shows the results of the whole-organism ELISA. All four IgY preparations demonstrated significant levels of activity, to a dilution of 1:62,500 or greater against both of the immunizing organism strains. Therefore, antibodies raised against one strain were highly cross-reactive with the other strain, and vice versa. The immunizing concentration of organisms did not have a significant effect on organism-specific IgY production, as both concentrations produced approximately equivalent responses. Therefore, the lower immunizing concentration of approximately 1.5 × 10⁸ organisms/hen is the preferred immunizing concentration of the two tested. The preimmune IgY preparation appeared to possess relatively low levels of *C. difficile*-reactive activity to a dilution of 1:500, probably due to prior exposure of the animals to environmental clostridia.

An initial whole-organism ELISA was performed using IgY preparations made from single CD 43594, #1 and CD 43596, #1 eggs collected around day 50 (data not shown). Specific titers were found to be 5 to 10-fold lower than those reported in Table 5. These results demonstrate that it is possible to begin immunizing hens prior to the time that they begin to lay eggs, and to obtain high titer specific IgY from the first eggs that are laid. In other words, it is not necessary to wait for the hens to begin laying before the immunization schedule is started.

TABLE 5

Results Of The Anti-C. difficile Whole-Organism ELISA

IgY Preparation	Dilution Of IgY Prep	43594-Coated Wells	43596-Coated Wells
	<u> </u>	433X4-Coated Wells	~43330-Coaled Wells
	1:500	1.746	1.801
	1:2,500	1.092	1.670
CD 43594, #1	1:12,500	0.202	0.812
, , , , , ,	1:62,500	0.136	0.179
	1:312,500	0.012	0.080
	1:1,562,500	0.002	0.020
	1:500	1.780	1.771
i i	1:2,500	1.025	1.078
CD 43594, #7	1:12,500	0.188	0.382
CD 43334, #7	1:62,500	0.052	0.132
	1:312,500	0.022	0.043
	1:1,562,500	0.005	0.024
	1:500	1.526	1.790
	1:2,500	0.832	1.477
CD 43596, #1	1:12,500	0.247	0.452
CD 43390, #1	1:62,500	0.050	0.242
	1:312,500	0.010	0.067
	1:1,562,500	0.000	0.036
	1:500	1.702	1.505
`	1:2,500	0.706	0.866
CD 43596, #7	1:12,500	0.250	0.282
CD 43390, #7	1:62,500	0.039	0.078
	1:312,500	0.002	0.017
	1:1,562,500	0.000	0.010
	1:500	0.142	0.309
j	1:2,500	0.032	0.077
Preimmune IgY	1:12,500	0.006	0.024
r remaindine ig r	1:62,500	0.002	0.012
	1:312,500	0.004	0.010
	1:1,562,500	0.002	0.014

EXAMPLE 2

Treatment Of C. difficile Infection With Anti-C. difficile Antibody

In order to determine whether the immune IgY antibodies raised against whole C. difficile organisms were capable of inhibiting the infection of hamsters by C. difficile, hamsters infected by these bacteria were utilized. [Lyerly et al., Infect. Immun., 59:2215-2218 (1991).] This example involved: (a) determination of the

lethal dose of C. difficile organisms; and (b) treatment of infected animals with immune antibody or control antibody in nutritional solution.

a) Determination Of The Lethal Dose Of C. difficile Organisms

Determination of the lethal dose of *C. difficile* organisms was carried out according to the model described by D.M. Lyerly et al., Infect. Immun., 59:2215-2218 (1991). *C. difficile* strain ATCC 43596 (serogroup C, ATCC) was plated on BHI agar and grown anaerobically (BBL Gas Pak 100 system) at 37°C for 42 hours. Organisms

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were removed from the agar surface using a sterile dacron-tip swab and suspended in

sterile 0.9% NaCl solution to a density of 108 organisms/ml.

In order to determine the lethal dose of *C. difficile* in the presence of control antibody and nutritional formula, non-immune eggs were obtained from unimmunized hens and a 12% PEG preparation made as described in Example 1(c). This preparation was redissolved in one fourth the original yolk volume of vanilla flavor Ensure®.

Starting on day one, groups of female Golden Syrian hamsters (Harlan Sprague Dawley), 8-9 weeks old and weighing approximately 100 gm, were orally administered 1 ml of the preimmune/Ensure® formula at time zero, 2 hours, 6 hours, and 10 hours. At 1 hour, animals were orally administered 3.0 mg clindamycin HCl (Sigma) in 1 ml of water. This drug predisposes hamsters to *C. difficile* infection by altering the normal intestinal flora. On day two, the animals were given 1 ml of the preimmune IgY/Ensure® formula at time zero, 2 hours, 6 hours, and 10 hours. At 1 hour on day two, different groups of animals were inoculated orally with saline (control), or 10², 10⁴, 10⁶, or 10⁸ *C. difficile* organisms in 1 ml of saline. From days 3-12, animals were given 1 ml of the preimmune IgY/Ensure® formula three times daily and observed for the onset of diarrhea and death. Each animal was housed in an individual cage and was offered food and water *ad libitum*.

Administration of 10⁶ - 10⁸ organisms resulted in death in 3-4 days while the lower doses of 10² - 10⁴ organisms caused death in 5 days. Cecal swabs taken from dead animals indicated the presence of *C. difficile*. Given the effectiveness of the 10² dose, this number of organisms was chosen for the following experiment to see if hyperimmune anti-*C. difficile* antibody could block infection.

b) Treatment Of Infected Animals With Immune Antibody Or Control Antibody In Nutritional Formula

The experiment in (a) was repeated using three groups of seven hamsters each. Group A received no clindamycin or *C. difficile* and was the survival control. Group B received clindamycin, 10^2 *C. difficile* organisms and preimmune IgY on the same schedule as the animals in (a) above. Group C received clindamycin, 10^2 *C. difficile* organisms, and hyperimmune anti-*C. difficile* IgY on the same schedule as Group B. The anti-*C. difficile* IgY was prepared as described in Example 1 except that the 12% PEG preparation was dissolved in one fourth the original yolk volume of Ensure®.

All animals were observed for the onset of diarrhea or other disease symptoms and death. Each animal was housed in an individual cage and was offered food and water *ad libitum*. The results are shown in Table 6.

TABLE 6

The Effect Of Oral Feeding Of Hyperimmune IgY Antibody on C. difficile Infection

2 .	Animal Group	Time To Diarrhea	Time To Death
A	pre-immune IgY only	no diarrhea	no deaths
В	Clindamycin, C. difficile, preimmune IgY	30 hrs.	49 hrs.
С	Clindamycin, C. difficile, immune IgY	33 hrs.	56 hrs.

25 * Mean of seven animals.

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Hamsters in the control group A did not develop diarrhea and remained healthy during the experimental period. Hamsters in groups B and C developed diarrheal disease. Anti-C. difficile IgY did not protect the animals from diarrhea or death, all

animals succumbed in the same time interval as the animals treated with preimmune IgY. Thus, while immunization with whole organisms apparently can improve sublethal symptoms with particular bacteria (see U.S. Patent No. 5,080,895 to H. Tokoro), such an approach does not prove to be productive to protect against the lethal effects of *C. difficile*.

EXAMPLE 3

Production of C. botulinum Type A Antitoxin in Hens

In order to determine whether antibodies could be raised against the toxin produced by clostridial pathogens, which would be effective in treating clostridial diseases, antitoxin to *C. botulinum* type A toxin was produced. This example involves: (a) toxin modification; (b) immunization; (c) antitoxin collection; (d) antigenicity assessment; and (e) assay of antitoxin titer.

a) Toxin Modification

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C. botulinum type A toxoid was obtained from B. R. DasGupta. From this, the active type A neurotoxin (M.W. approximately 150 kD) was purified to greater than 99% purity, according to published methods. [B.R. DasGupta & V. Sathyamoorthy, Toxicon, 22:415 (1984).] The neurotoxin was detoxified with formaldehyde according to published methods. [B.R. Singh & B.R. DasGupta, Toxicon, 27:403 (1989).]

b) Immunization

C. botulinum toxoid for immunization was dissolved in PBS (1 mg/ml) and was emulsified with an approximately equal volume of CFA (GIBCO) for initial immunization or IFA for booster immunization. On day zero, two white leghorn hens, obtained from local breeders, were each injected at multiple sites (intramuscular and subcutaneous) with 1 ml inactivated toxoid emulsified in 1 ml CFA. Subsequent booster immunizations were made according to the following schedule for day of

injection and toxoid amount: days 14 and 21 - 0.5 mg; day 171 - 0.75 mg; days 394, 401, 409 - 0.25 mg. One hen received an additional booster of 0.150 mg on day 544.

c) Antitoxin Collection

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Total yolk immunoglobulin (IgY) was extracted as described in Example 1(c) and the IgY pellet was dissolved in the original yolk volume of PBS with thimerosal.

d) Antigenicity Assessment

Eggs were collected from day 409 through day 423 to assess whether the toxoid was sufficiently immunogenic to raise antibody. Eggs from the two hens were pooled and antibody was collected as described in the standard PEG protocol. [Example 1(c).] Antigenicity of the botulinal toxin was assessed on Western blots. The 150 kD detoxified type A neurotoxin and unmodified, toxic, 300 kD botulinal type A complex (toxin used for intragastric route administration for animal gut neutralization experiments; see Example 6) were separated on a SDS-polyacrylamide reducing gel. The Western blot technique was performed according to the method of Towbin. [H. Towbin et al., Proc. Natl. Acad. Sci. USA, 76:4350 (1979).] Ten µg samples of C. botulinum complex and toxoid were dissolved in SDS reducing sample buffer (1% SDS, 0.5% 2-mercaptoethanol, 50 mM Tris, pH 6.8, 10% glycerol, 0.025% w/v bromphenol blue, 10% β -mercaptoethanol), heated at 95°C for 10 min and separated on a 1 mm thick 5% SDS-polyacrylamide gel. [K. Weber and M. Osborn,"Proteins and Sodium Dodecyl Sulfate: Molecular Weight Determination on Polyacrylamide Gels and Related Procedures," in The Proteins, 3d Edition (H. Neurath & R.L. Hill, eds), pp. 179-223, (Academic Press, NY, 1975).] Part of the gel was cut off and the proteins were stained with Coomassie Blue. The proteins in the remainder of the gel were transferred to nitrocellulose using the Milliblot-SDE electroblotting system (Millipore) according to manufacturer's directions. The nitrocellulose was temporarily stained with 10% Ponceau S [S.B. Carroll and A. Laughon, "Production and Purification of Polyclonal Antibodies to the Foreign Segment of \(\beta -

galactosidase Fusion Proteins," in DNA Cloning: A Practical Approach, Vol.III, (D. Glover, ed.), pp. 89-111, IRL Press, Oxford, (1987)] to visualize the lanes, then destained by running a gentle stream of distilled water over the blot for several minutes. The nitrocellulose was immersed in PBS containing 3% BSA overnight at 4°C to block any remaining protein binding sites.

The blot was cut into strips and each strip was incubated with the appropriate primary antibody. The avian anti-C. botulinum antibodies [described in (c)] and pre-immune chicken antibody (as control) were diluted 1:125 in PBS containing 1 mg/ml BSA for 2 hours at room temperature. The blots were washed with two changes each of large volumes of PBS, BBS-Tween and PBS, successively (10 min/wash). Goat anti-chicken IgG alkaline phosphatase conjugated secondary antibody (Fisher Biotech) was diluted 1:500 in PBS containing 1 mg/ml BSA and incubated with the blot for 2 hours at room temperature. The blots were washed with two changes each of large volumes of PBS and BBS-Tween, followed by one change of PBS and 0.1 M Tris-HCl, pH 9.5. Blots were developed in freshly prepared alkaline phosphatase substrate buffer (100 µg/ml nitroblue tetrazolium (Sigma), 50 µg/ml 5-bromo-4-chloro-3-indolyl phosphate (Sigma), 5 mM MgCl₂ in 50 mM Na₂CO₃, pH 9.5).

The Western blots are shown in Figure 1. The anti-C. botulinum IgY reacted to the toxoid to give a broad immunoreactive band at about 145-150 kD on the reducing gel. This toxoid is refractive to disulfide cleavage by reducing agents due to formalin crosslinking. The immune IgY reacted with the active toxin complex, a 97 kD C. botulinum type A heavy chain and a 53 kD light chain. The preimmune IgY was unreactive to the C. botulinum complex or toxoid in the Western blot.

e) Antitoxin Antibody Titer

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The IgY antibody titer to *C. botulinum* type A toxoid of eggs harvested between day 409 and 423 was evaluated by ELISA, prepared as follows. Ninety-six-well Falcon Pro-bind plates were coated overnight at 4°C with 100 µl/well toxoid [B.R. Singh & B.R. Das Gupta, Toxicon 27:403 (1989)] at 2.5 µg/ml in PBS, pH 7.5

containing 0.005% thimerosal. The following day the wells were blocked with PBS containing 1% BSA for 1 hour at 37°C. The IgY from immune or preimmune eggs was diluted in PBS containing 1% BSA and 0.05% Tween 20 and the plates were incubated for 1 hour at 37°C. The plates were washed three times with PBS containing 0.05% Tween 20 and three times with PBS alone. Alkaline phosphatase-conjugated goat-anti-chicken IgG (Fisher Biotech) was diluted 1:750 in PBS containing 1% BSA and 0.05% Tween 20, added to the plates, and incubated 1 hour at 37°C. The plates were washed as before, and p-nitrophenyl phosphate (Sigma) at 1 mg/ml in 0.05 M Na₂CO₃, pH 9.5, 10 mM MgCl₂ was added.

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The results are shown in Figure 2. Chickens immunized with the toxoid generated high titers of antibody to the immunogen. Importantly, eggs from both immunized hens had significant anti-immunogen antibody titers as compared to preimmune control eggs. The anti-C. botulinum IgY possessed significant activity, to a dilution of 1:93,750 or greater.

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EXAMPLE 4

Preparation Of Avian Egg Yolk Immunoglobulin In An Orally Administrable Form

In order to administer avian IgY antibodies orally to experimental mice, an effective delivery formula for the IgY had to be determined. The concern was that if the crude IgY was dissolved in PBS, the saline in PBS would dehydrate the mice, which might prove harmful over the duration of the study. Therefore, alternative methods of oral administration of IgY were tested. The example involved: (a) isolation of immune IgY; (b) solubilization of IgY in water or PBS, including subsequent dialysis of the IgY-PBS solution with water to eliminate or reduce the salts (salt and phosphate) in the buffer; and (c) comparison of the quantity and activity of recovered IgY by absorbance at 280 nm and PAGE, and enzyme-linked immunoassay (ELISA).

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a) Isolation Of Immune IgY

In order to investigate the most effective delivery formula for IgY, we used IgY which was raised against *Crotalus durissus terrificus* venom. Three eggs were collected from hens immunized with the *C. durissus terrificus* venom and IgY was extracted from the yolks using the modified Polson procedure described by Thalley and Carroll [Bio/Technology, 8:934-938 (1990)] as described in Example 1(c).

The egg yolks were separated from the whites, pooled, and blended with four volumes of PBS. Powdered PEG 8000 was added to a concentration of 3.5%. The mixture was centrifuged at 10,000 rpm for 10 minutes to pellet the precipitated protein, and the supernatant was filtered through cheesecloth to remove the lipid layer. Powdered PEG 8000 was added to the supernatant to bring the final PEG concentration to 12% (assuming a PEG concentration of 3.5% in the supernatant). The 12% PEG/IgY mixture was divided into two equal volumes and centrifuged to pellet the IgY.

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b) Solubilization Of The IgY In Water Or PBS

One pellet was resuspended in 1/2 the original yolk volume of PBS, and the other pellet was resuspended in 1/2 the original yolk volume of water. The pellets were then centrifuged to remove any particles or insoluble material. The IgY in PBS solution dissolved readily but the fraction resuspended in water remained cloudy.

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In order to satisfy anticipated sterility requirements for orally administered antibodies, the antibody solution needs to be filter-sterilized (as an alternative to heat sterilization which would destroy the antibodies). The preparation of IgY resuspended in water was too cloudy to pass through either a 0.2 or 0.45 µm membrane filter, so 10 ml of the PBS resuspended fraction was dialyzed overnight at room temperature against 250 ml of water. The following morning the dialysis chamber was emptied and refilled with 250 ml of fresh H₂O for a second dialysis. Thereafter, the yields of soluble antibody were determined at OD₂₈₀ and are compared in Table 7.

TABLE 7

Dependence Of IgY Yield On Solvents

Fraction Abs	orbance Of 1:10 Dilution At 280	nm Percent Recovery
PBS dissolved	1.149	100%
H₂O dissolved	0.706	61%
PBS dissolved/H ₂ O dialyzed	0.885	77%

Resuspending the pellets in PBS followed by dialysis against water recovered more antibody than directly resuspending the pellets in water (77% versus 61%). Equivalent volumes of the IgY preparation in PBS or water were compared by PAGE, and these results were in accordance with the absorbance values (data not shown).

c) Activity Of IgY Prepared With Different Solvents

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An ELISA was performed to compare the binding activity of the IgY extracted by each procedure described above. *C. durissus terrificus* (*C.d.t.*) venom at 2.5 µg/ml in PBS was used to coat each well of a 96-well microtiter plate. The remaining protein binding sites were blocked with PBS containing 5 mg/ml BSA. Primary antibody dilutions (in PBS containing 1 mg/ml BSA) were added in duplicate. After 2 hours of incubation at room temperature, the unbound primary antibodies were removed by washing the wells with PBS, BBS-Tween, and PBS. The species specific secondary antibody (goat anti-chicken immunoglobulin alkaline-phosphatase conjugate (Sigma) was diluted 1:750 in PBS containing 1 mg/ml BSA and added to each well of the microtiter plate. After 2 hours of incubation at room temperature, the unbound secondary antibody was removed by washing the plate as before, and freshly prepared alkaline phosphatase substrate (Sigma) at 1 mg/ml in 50 mM Na₂CO₃, 10 mM MgCl₂, pH 9.5 was added to each well. The color development was measured on a Dynatech MR 700 microplate reader using a 412 nm filter. The results are shown in Table 8.

The binding assay results parallel the recovery values in Table 7, with PBS-dissolved IgY showing slightly more activity than the PBS-dissolved/H₂O dialyzed

antibody. The water-dissolved antibody had considerably less binding activity than the other preparations.

EXAMPLE 5

Survival Of Antibody Activity After Passage Through The Gastrointestinal Tract

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In order to determine the feasibility of oral administration of antibody, it was of interest to determine whether orally administered IgY survived passage through the gastrointestinal tract. The example involved: (a) oral administration of specific immune antibody mixed with a nutritional formula; and (b) assay of antibody activity extracted from feces.

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TABLE 8

Antigen-Binding Activity Of IgY Prepared With Different Solvents

Dilution	Preimmune	PBS Dissolved	H₂O Dissolved	PBS/H ₂ O
1:500	0.005	1.748	1.577	1.742
1:2,500	0.004	0.644	0.349	0.606
1:12,500	0.001	0.144	0.054	0.090
1:62,500	0.001	0.025	0.007	0.016
1:312,500	0.010	0.000	0.000	0.002

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a) Oral Administration Of Antibody

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The IgY preparations used in this example are the same PBS-dissolved/H₂O dialyzed antivenom materials obtained in Example 4 above, mixed with an equal volume of Enfamil®. Two mice were used in this experiment, each receiving a different diet as follows:

- 1) water and food as usual;
- 2) immune IgY preparation dialyzed against water and mixed 1:1 with 25 Enfamil®. (The mice were given the corresponding mixture as their only source of food and water).

b) Antibody Activity After Ingestion

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After both mice had ingested their respective fluids, each tube was refilled with approximately 10 ml of the appropriate fluid first thing in the morning. By midmorning there was about 4 to 5 ml of liquid left in each tube. At this point stool samples were collected from each mouse, weighed, and dissolved in approximately 500 µl PBS per 100 mg stool sample. One hundred and sixty mg of control stools (no antibody) and 99 mg of experimental stools (specific antibody) in 1.5 ml microfuge tubes were dissolved in 800 and 500 µl PBS, respectively. The samples were heated at 37°C for 10 minutes and vortexed vigorously. The experimental stools were also broken up with a narrow spatula. Each sample was centrifuged for 5 minutes in a microfuge and the supernatants, presumably containing the antibody extracts, were collected. The pellets were saved at 2-8°C in case future extracts were needed. Because the supernatants were tinted, they were diluted five-fold in PBS containing 1 mg/ml BSA for the initial dilution in the enzyme immunoassay (ELISA). The primary extracts were then diluted five-fold serially from this initial dilution. The volume of primary extract added to each well was 190 µl. The ELISA was performed exactly as described in Example 4.

TABLE 9

Specific Antibody Activity After Passage Through The Gastrointestinal Tract

Dilution	Preimmune IgY	Control Fecal Extract	EXP. Fecal Extract
1:5	<0	0.000	0.032
1:25	0.016	<0	0.016
1:125	<0	<0 .	0.009
1:625	<0	0.003	0.001
1:3125	<0	<0	0.000

There was some active antibody in the fecal extract from the mouse given the specific antibody in Enfamil® formula, but it was present at a very low level. Since the samples were assayed at an initial 1:5 dilution, the binding observed could have been higher with less dilute samples. Consequently, the mice were allowed to

continue ingesting either regular food and water or the specific IgY in Enfamil® formula, as appropriate, so the assay could be repeated. Another ELISA plate was coated overnight with 5 µg/ml of C.d.t. venom in PBS.

The following morning the ELISA plate was blocked with 5 mg/ml BSA, and the fecal samples were extracted as before, except that instead of heating the extracts at 37°C, the samples were kept on ice to limit proteolysis. The samples were assayed undiluted initially, and in 5X serial dilutions thereafter. Otherwise the assay was carried out as before.

TABLE 10
Specific Antibody Survives Passage Through The Gastrointestinal Tract

Dilution	Preimmune IgY	Control Extract	Exp. Extract
undiluted	0.003	<0	0.379
1:5	<0	<0	0.071
1:25	0.000	<0	0.027
1:125	0.003	<0	0.017
1:625	0.000	<0	0.008
1:3125	0.002	<0	0.002

The experiment confirmed the previous results, with the antibody activity markedly higher. The control fecal extract showed no anti-C.d.t. activity, even undiluted, while the fecal extract from the anti-C.d.t. IgY/Enfamil®-fed mouse showed considerable anti-C.d.t. activity. This experiment (and the previous experiment) clearly demonstrate that active IgY antibody survives passage through the mouse digestive tract, a finding with favorable implications for the success of IgY antibodies administered orally as a therapeutic or prophylactic.

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EXAMPLE 6

In Vivo Neutralization Of Type C. botulinum Type A Neurotoxin By Avian Antitoxin Antibody

This example demonstrated the ability of PEG-purified antitoxin, collected as described in Example 3, to neutralize the lethal effect of C. botulinum neurotoxin type A in mice. To determine the oral lethal dose (LD₁₀₀) of toxin A, groups of BALB/c mice were given different doses of toxin per unit body weight (average body weight of 24 grams). For oral administration, toxin A complex, which contains the neurotoxin associated with other non-toxin proteins was used. This complex is markedly more toxic than purified neurotoxin when given by the oral route. [I. Ohishi et al., Infect. Immun., 16:106 (1977).] C. botulinum toxin type A complex, obtained from Eric Johnson (University Of Wisconsin, Madison) was 250 μg/ml in 50 mM sodium citrate, pH 5.5, specific toxicity 3×10^7 mouse LD₅₀/mg with parenteral administration. Approximately 40-50 ng/gm body weight was usually fatal within 48 hours in mice maintained on conventional food and water. When mice were given a diet ad libitum of only Enfamil® the concentration needed to produce lethality was approximately 2.5 times higher (125 ng/gm body weight). Botulinal toxin concentrations of approximately 200 ng/gm body weight were fatal in mice fed Enfamil® containing preimmune IgY (resuspended in Enfamil® at the original yolk volume).

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The oral LD₁₀₀ of *C. botulinum* toxin was also determined in mice that received known amounts of a mixture of preimmune IgY-Ensure® delivered orally through feeding needles. Using a 22 gauge feeding needle, mice were given 250 µl each of a preimmune IgY-Ensure® mixture (preimmune IgY dissolved in 1/4 original yolk volume) 1 hour before and 1/2 hour and 5 hours after administering botulinal toxin. Toxin concentrations given orally ranged from approximately 12 to 312 ng/gm body weight (0.3 to 7.5 µg per mouse). Botulinal toxin complex concentration of approximately 40 ng/gm body weight (1 µg per mouse) was lethal in all mice in less than 36 hours.

Two groups of BALB/c mice, 10 per group, were each given orally a single dose of 1 µg each of botulinal toxin complex in 100 µl of 50 mM sodium citrate pH 5.5. The mice received 250 µl treatments of a mixture of either preimmune or immune IgY in Ensure® (1/4 original yolk volume) 1 hour before and 1/2 hour, 4 hours, and 8 hours after botulinal toxin administration. The mice received three treatments per day for two more days. The mice were observed for 96 hours. The survival and mortality are shown in Table 11.

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TABLE 11
Neutralization Of Botulinal Toxin A In Vivo

Toxin Dose ng/gm	Antibody Type	Number Of Mice Alive	Number Of Mice Dead
41.6	non-immune	0	10
41.6	anti-botulinal toxin	10	0

All mice treated with the preimmune IgY-Ensure® mixture died within 46 hours post-toxin administration. The average time of death in the mice was 32 hours post toxin administration. Treatments of preimmune IgY-Ensure® mixture did not continue beyond 24 hours due to extensive paralysis of the mouth in mice of this group. In contrast, all ten mice treated with the immune anti-botulinal toxin IgY-Ensure® mixture survived past 96 hours. Only 4 mice in this group exhibited symptoms of botulism toxicity (two mice about 2 days after and two mice 4 days after toxin administration). These mice eventually died 5 and 6 days later. Six of the mice in this immune group displayed no adverse effects to the toxin and remained alive and healthy long term. Thus, the avian anti-botulinal toxin antibody demonstrated very good protection from the lethal effects of the toxin in the experimental mice.

EXAMPLE 7

Production Of An Avian Antitoxin Against Clostridium difficile Toxin A

Toxin A is a potent cytotoxin secreted by pathogenic strains of C. difficile, that plays a direct role in damaging gastrointestinal tissues. In more severe cases of C. difficile intoxication, pseudomembranous colitis can develop which may be fatal. This would be prevented by neutralizing the effects of this toxin in the gastrointestinal tract. As a first step, antibodies were produced against a portion of the toxin. The example involved: (a) conjugation of a synthetic peptide of toxin A to bovine serum albumin; (b) immunization of hens with the peptide-BSA conjugate; and (c) detection of antitoxin peptide antibodies by ELISA.

a) Conjugation Of A Synthetic Peptide Of Toxin A To Bovine Serum Albumin

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The synthetic peptide CQTIDGKKYYFN-NH₂ (SEQ ID NO:82) was prepared commercially (Multiple Peptide Systems, San Diego, CA) and validated to be >80% pure by high-pressure liquid chromatography. The eleven amino acids following the cysteine residue represent a consensus sequence of a repeated amino acid sequence found in Toxin A. [Wren et al., Infect. Immun., 59:3151-3155 (1991).] The cysteine was added to facilitate conjugation to carrier protein.

In order to prepare the carrier for conjugation, BSA (Sigma) was dissolved in 0.01 M NaPO₄, pH 7.0 to a final concentration of 20 mg/ml and n-maleimidobenzoyl-N-hydroxysuccinimide ester (MBS; Pierce) was dissolved in N,N-dimethyl formamide to a concentration of 5 mg/ml. MBS solution, 0.51 ml, was added to 3.25 ml of the BSA solution and incubated for 30 minutes at room temperature with stirring every 5 minutes. The MBS-activated BSA was then purified by chromatography on a Bio-Gel P-10 column (Bio-Rad; 40 ml bed volume) equilibrated with 50 mM NaPO₄, pH 7.0 buffer. Peak fractions were pooled (6.0 ml).

Lyophilized toxin A peptide (20 mg) was added to the activated BSA mixture, stirred until the peptide dissolved and incubated 3 hours at room temperature. Within 20 minutes, the reaction mixture became cloudy and precipitates formed. After 3 hours, the reaction mixture was centrifuged at 10,000 × g for 10 min and the supernatant analyzed for protein content. No significant protein could be detected at 280 nm. The conjugate precipitate was washed three times with PBS and stored at 4°C. A second conjugation was performed with 15 mg of activated BSA and 5 mg of peptide and the conjugates pooled and suspended at a peptide concentration of 10 mg/ml in 10 mM NaPO₄, pH 7.2.

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b) Immunization Of Hens With Peptide Conjugate

Two hens were each initially immunized on day zero by injection into two subcutaneous and two intramuscular sites with 1 mg of peptide conjugate that was emulsified in CFA (GIBCO). The hens were boosted on day 14 and day 21 with 1 mg of peptide conjugate emulsified in IFA (GIBCO).

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c) Detection Of Antitoxin Peptide Antibodies By ELISA

IgY was purified from two eggs obtained before immunization (pre-immune) and two eggs obtained 31 and 32 days after the initial immunization using PEG fractionation as described in Example 1.

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Wells of a 96-well microtiter plate (Falcon Pro-Bind Assay Plate) were coated overnight at 4°C with 100 µg/ml solution of the toxin A synthetic peptide in PBS, pH 7.2 prepared by dissolving 1 mg of the peptide in 1.0 ml of H₂O and dilution of PBS. The pre-immune and immune IgY preparations were diluted in a five-fold series in a buffer containing 1% PEG 8000 and 0.1% Tween-20 (v/v) in PBS, pH 7.2. The wells were blocked for 2 hours at room temperature with 150 µl of a solution containing 5% (v/v) Carnation® nonfat dry milk and 1% PEG 8000 in PBS, pH 7.2. After incubation for 2 hours at room temperature, the wells were washed, secondary rabbit anti-chicken IgG-alkaline phosphatase (1:750) added, the wells washed again and the

color development obtained as described in Example 1. The results are shown in Table 12.

TABLE 12

Reactivity Of IgY With Toxin Peptide

Dilution Of PEG Prep	Absorbance At 410 nm Preimmune Immune Anti-Peptide		
1:100	0.013	0.253	
1:500	0.004	0.039	
1:2500	0.004	0.005	

Clearly, the immune antibodies contain titers against this repeated epitope of toxin A.

EXAMPLE 8

Production Of Avian Antitoxins Against Clostridium difficile Native Toxins A And B

To determine whether avian antibodies are effective for the neutralization of C. difficile toxins, hens were immunized using native C. difficile toxins A and B. The resulting egg yolk antibodies were then extracted and assessed for their ability to neutralize toxins A and B in vitro. The Example involved (a) preparation of the toxin immunogens, (b) immunization, (c) purification of the antitoxins, and (d) assay of toxin neutralization activity.

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a) Preparation Of The Toxin Immunogens

Both C. difficile native toxins A and B, and C. difficile toxoids, prepared by the treatment of the native toxins with formaldehyde, were employed as immunogens.

C. difficile toxoids A and B were prepared by a procedure which was modified from published methods (Ehrich et al., Infect. Immun. 28:1041 (1980). Separate solutions (in PBS) of native C. difficile toxin A and toxin B (Tech Lab) were each adjusted to a

concentration of 0.20 mg/ml, and formaldehyde was added to a final concentration of 0.4%. The toxin/formaldehyde solutions were then incubated at 37°C for 40 hrs. Free formaldehyde was then removed from the resulting toxoid solutions by dialysis against PBS at 4°C. In previously published reports, this dialysis step was not performed. Therefore, free formaldehyde must have been present in their toxoid preparations. The toxoid solutions were concentrated, using a Centriprep concentrator unit (Amicon), to a final toxoid concentration of 4.0 mg/ml. The two resulting preparations were designated as toxoid A and toxoid B.

C. difficile native toxins were prepared by concentrating stock solutions of toxin A and toxin B (Tech Lab, Inc), using Centriprep concentrator units (Amicon), to a final concentration of 4.0 mg/ml.

b) Immunization

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The first two immunizations were performed using the toxoid A and toxoid B immunogens described above. A total of 3 different immunization combinations were employed. For the first immunization group, 0.2 ml of toxoid A was emulsified in an equal volume of Titer Max adjuvant (CytRx). Titer Max was used in order to conserve the amount of immunogen used, and to simplify the immunization procedure. This immunization group was designated "CTA." For the second immunization group, 0.1 ml of toxoid B was emulsified in an equal volume of Titer Max adjuvant. This group was designated "CTB." For the third immunization group, 0.2 ml of toxoid A was first mixed with 0.2 ml of toxoid B, and the resulting mixture was emulsified in 0.4 ml of Titer Max adjuvant. This group was designated "CTAB." In this way, three separate immunogen emulsions were prepared, with each emulsion containing a final concentration of 2.0 mg/ml of toxoid A (CTA) or toxoid B (CTB) or a mixture of 2.0 mg/ml toxoid A and 2.0 mg/ml toxoid B (CTAB).

On day 0, White Leghorn hens, obtained from a local breeder, were immunized as follows: Group CTA. Four hens were immunized, with each hen receiving 200µg of toxoid A, via two intramuscular (I.M.) injections of 50µl of CTA emulsion in the breast area. Group CTB. One hen was immunized with 200µg of toxoid B, via two

I.M. injections of 50µl of CTB emulsion in the breast area. Group CTAB. Four hens were immunized, with each hen receiving a mixture containing 200µg of toxoid A and 200µg of toxoid B, via two I.M. injections of 100µl of CTAB emulsion in the breast area. The second immunization was performed 5 weeks later, on day 35, exactly as described for the first immunization above.

In order to determine whether hens previously immunized with *C. difficile* toxoids could tolerate subsequent booster immunizations using native toxins, a single hen from group CTAB was immunized for a third time, this time using a mixture of the native toxin A and native toxin B described in section (a) above (these toxins were not formaldehyde-treated, and were used in their active form). This was done in order to increase the amount (titer) and affinity of specific antitoxin antibody produced by the hen over that achieved by immunizing with toxoids only. On day 62, 0.1 ml of a toxin mixture was prepared which contained 200µg of native toxin A and 200µg of native toxin B. This toxin mixture was then emulsified in 0.1 ml of Titer Max adjuvant. A single CTAB hen was then immunized with the resulting immunogen emulsion, via two I.M. injections of 100µl each, into the breast area. This hen was marked with a wing band, and observed for adverse effects for a period of approximately 1 week, after which time the hen appeared to be in good health.

Because the CTAB hen described above tolerated the booster immunization with native toxins A and B with no adverse effects, it was decided to boost the remaining hens with native toxin as well. On day 70, booster immunizations were performed as follows: Group CTA. A 0.2 ml volume of the 4 mg/ml native toxin A solution was emulsified in an equal volume of Titer Max adjuvant. Each of the 4 hens was then immunized with 200µg of native toxin A, as described for the toxoid A immunizations above. Group CTB. A 50µl volume of the 4 mg/ml native toxin B solution was emulsified in an equal volume of Titer Max adjuvant. The hen was then immunized with 200µg of native toxin B, as described for the toxoid B immunizations above. Group CTAB. A 0.15 ml volume of the 4 mg/ml native toxin A solution was first mixed with a 0.15 ml volume the 4 mg/ml native toxin B solution. The resulting toxin mixture was then emulsified in 0.3 ml of Titer Max adjuvant. The 3 remaining

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hens (the hen with the wing band was not immunized this time) were then immunized with 200µg of native toxin A and 200µg of native toxin B as described for the toxoid A+ toxoid B immunizations (CTAB) above. On day 85, all hens received a second booster immunization using native toxins, done exactly as described for the first boost with native toxins above.

All hens tolerated both booster immunizations with native toxins with no adverse effects. As previous literature references describe the use of formaldehydetreated toxoids, this is apparently the first time that any immunizations have been performed using native C. difficile toxins.

c) Purification Of Antitoxins

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Eggs were collected from the hen in group CTB 10-12 days following the second immunization with toxoid (day 35 immunization described in section (b) above), and from the hens in groups CTA and CTAB 20-21 days following the second immunization with toxoid. To be used as a pre-immune (negative) control, eggs were also collected from unimmunized hens from the same flock. Egg yolk immunoglobulin (IgY) was extracted from the 4 groups of eggs as described in Example 1 (c), and the final IgY pellets were solubilized in the original yolk volume of PBS without thimerosal. Importantly, thimerosal was excluded because it would have been toxic to the CHO cells used in the toxin neutralization assays described in section (d) below.

d) Assay Of Toxin Neutralization Activity

The toxin neutralization activity of the IgY solutions prepared in section (c) above was determined using an assay system that was modified from published methods. [Ehrich et al., Infect. Immun. 28:1041-1043 (1992); and McGee et al. Microb. Path. 12:333-341 (1992).] As additional controls, affinity-purified goat anti-C. difficile toxin A (Tech Lab) and affinity-purified goat anti-C. difficile toxin B (Tech Lab) were also assayed for toxin neutralization activity.

The IgY solutions and goat antibodies were serially diluted using F 12 medium (GIBCO) which was supplemented with 2% FCS (GIBCO)(this solution will be referred to as "medium" for the remainder of this Example). The resulting antibody solutions were then mixed with a standardized concentration of either native C. difficile toxin A (Tech Lab), or native C. difficile toxin B (Tech Lab), at the concentrations indicated below. Following incubation at 37°C for 60 min., 100µl volumes of the toxin + antibody mixtures were added to the wells of 96-well microtiter plates (Falcon Microtest III) which contained 2.5 x 10⁴ Chinese Hamster Ovary (CHO) cells per well (the CHO cells were plated on the previous day to allow them to adhere to the plate wells). The final concentration of toxin, or dilution of antibody indicated below refers to the final test concentration of each reagent present in the respective microtiter plate wells. Toxin reference wells were prepared which contained CHO cells and toxin A or toxin B at the same concentration used for the toxin plus antibody mixtures (these wells contained no antibody). Separate control wells were also prepared which contained CHO cells and medium only. The assay plates were then incubated for 18-24 hrs. in a 37°C, humidified, 5% CO2 incubator. On the following day, the remaining adherent (viable) cells in the plate wells were stained using 0.2% crystal violet (Mallinckrodt) dissolved in 2% ethanol, for 10 min. Excess stain was then removed by rinsing with water, and the stained cells were solubilized by adding 100µl of 1% SDS (dissolved in water) to each well. The absorbance of each well was then measured at 570 nm, and the percent cytotoxicity of each test sample or mixture was calculated using the following formula:

% CHO Cell Cytotoxicity =
$$[1 - (\frac{Abs. Sample}{Abs. Control})] X 100$$

Unlike previous reports which quantitate results visually by counting cell rounding by microscopy, this Example utilized spectrophotometric methods to quantitate the *C. difficile* toxin bioassay. In order to determine the toxin A

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neutralizing activity of the CTA, CTAB, and pre-immune IgY preparations, as well as the affinity-purified goat antitoxin A control, dilutions of these antibodies were reacted against a 0.1µg/ml concentration of native toxin A (this is the approx. 50% cytotoxic dose of toxin A in this assay system). The results are shown in Figure 3.

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Complete neutralization of toxin A occurred with the CTA IgY (antitoxin A, above) at dilutions of 1:80 and lower, while significant neutralization occurred out to the 1:320 dilution. The CTAB IgY (antitoxin A + toxin B, above) demonstrated complete neutralization at the 1:320-1:160 and lower dilutions, and significant neutralization occurred out to the 1:1280 dilution. The commercially available affinity-purified goat antitoxin A did not completely neutralize toxin A at any of the dilutions tested, but demonstrated significant neutralization out to a dilution of 1:1,280. The preimmune IgY did not show any toxin A neutralizing activity at any of the concentrations tested. These results demonstrate that IgY purified from eggs laid by hens immunized with toxin A alone, or simultaneously with toxin A and toxin B, is an effective toxin A antitoxin.

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The toxin B neutralizing activity of the CTAB and pre-immune IgY preparations, and also the affinity-purified goat antitoxin B control was determined by reacting dilutions of these antibodies against a concentration of native toxin B of 0.1 ng/ml (approximately the 50% cytotoxic dose of toxin B in the assay system). The results are shown in Figure 4.

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Complete neutralization of toxin B occurred with the CTAB IgY (antitoxin A + toxin B, above) at the 1:40 and lower dilutions, and significant neutralization occurred out to the 1:320 dilution. The affinity-purified goat antitoxin B demonstrated complete neutralization at dilutions of 1:640 and lower, and significant neutralization occurred out to a dilution of 1:2,560. The preimmune IgY did not show any toxin B neutralizing activity at any of the concentrations tested. These results demonstrate that IgY purified from eggs laid by hens immunized simultaneously with toxin A and toxin B is an effective toxin B antitoxin.

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In a separate study, the toxin B neutralizing activity of CTB, CTAB, and preimmune IgY preparations was determined by reacting dilutions of these antibodies

against a native toxin B concentration of 0.1µg/ml (approximately 100% cytotoxic dose of toxin B in this assay system). The results are shown in Figure 5.

Significant neutralization of toxin B occurred with the CTB IgY (antitoxin B, above) at dilutions of 1:80 and lower, while the CTAB IgY (antitoxin A + toxin B, above) was found to have significant neutralizing activity at dilutions of 1:40 and lower. The preimmune IgY did not show any toxin B neutralizing activity at any of the concentrations tested. These results demonstrate that IgY purified from eggs laid by hens immunized with toxin B alone, or simultaneously with toxin A and toxin B, is an effective toxin B antitoxin.

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EXAMPLE 9

In vivo Protection Of Golden Syrian Hamsters From

C. difficile Disease By Avian Antitoxins Against C. difficile Toxins A And B

The most extensively used animal model to study *C. difficile* disease is the hamster. [Lyerly et al., Infect. Immun. 47:349-352 (1992).] Several other animal models for antibiotic-induced diarrhea exist, but none mimic the human form of the disease as closely as the hamster model. [R. Fekety, "Animal Models of Antibiotic-Induced Colitis," in O. Zak and M. Sande (eds.), Experimental Models in Antimicrobial Chemotherapy, Vol. 2, pp.61-72, (1986).] In this model, the animals are first predisposed to the disease by the oral administration of an antibiotic, such as clindamycin, which alters the population of normally-occurring gastrointestinal flora (Fekety, at 61-72). Following the oral challenge of these animals with viable C. difficile organisms, the hamsters develop cecitis, and hemorrhage, ulceration, and inflammation are evident in the intestinal mucosa. [Lyerly et al., Infect. Immun. 47:349-352 (1985).] The animals become lethargic, develop severe diarrhea, and a high percentage of them die from the disease. [Lyerly et al., Infect. Immun. 47:349-352 (1985).] This model is therefore ideally suited for the evaluation of therapeutic agents designed for the treatment or prophylaxis of C. difficile disease.

The ability of the avian C. difficile antitoxins, described in Example 1 above, to protect hamsters from C. difficile disease was evaluated using the Golden Syrian hamster model of C. difficile infection. The Example involved (a) preparation of the avian C. difficile antitoxins, (b) in vivo protection of hamsters from C. difficile disease by treatment with avian antitoxins, and (c) long-term survival of treated hamsters.

a) Preparation Of The Avian C. difficile Antitoxins

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Eggs were collected from hens in groups CTA and CTAB described in Example 1 (b) above. To be used as a pre-immune (negative) control, eggs were also purchased from a local supermarket. Egg yolk immunoglobulin (IgY) was extracted from the 3 groups of eggs as described in Example 1 (c), and the final IgY pellets were solubilized in one fourth the original yolk volume of Ensure® nutritional formula.

b) In vivo Protection Of Hamsters Against C. difficile Disease By Treatment With Avian Antitoxins

The avian C. difficile antitoxins prepared in section (a) above were evaluated 15 for their ability to protect hamsters from C. difficile disease using an animal model system which was modified from published procedures. [Fekety, at 61-72; Bortiello et al., J. Med. Microbiol., 24:53-64 (1987); Kim et al., Infect. Immun., 55:2984-2992 (1987); Borriello et al., J. Med. Microbiol., 25:191-196 (1988); Delmee and Avesani, 20 J. Med. Microbiol., 33:85-90 (1990); and Lyerly et al., Infect. Immun., 59:2215-2218 (1991).] For the study, three separate experimental groups were used, with each group consisting of 7 female Golden Syrian hamsters (Charles River), approximately 10 weeks old and weighing approximately 100 gms. each. The three groups were designated "CTA." "CTAB" and "Pre-immune." These designations corresponded to the antitoxin preparations with which the animals in each group were treated. Each 25 animal was housed in an individual cage, and was offered food and water ad libitum through the entire length of the study. On day 1, each animal was orally administered

1.0 ml of one of the three antitoxin preparations (prepared in section (a) above) at the following timepoints: 0 hrs., 4 hrs., and 8 hrs. On day 2, the day 1 treatment was repeated. On day 3, at the 0 hr. timepoint, each animal was again administered antitoxin, as described above. At 1 hr., each animal was orally administered 3.0 mg of clindamycin-HCl (Sigma) in 1 ml of water. This treatment predisposed the animals to infection with C. difficile. As a control for possible endogenous C. difficile colonization, an additional animal from the same shipment (untreated) was also administered 3.0 mg of clindamycin-HCl in the same manner. This clindamycin control animal was left untreated (and uninfected) for the remainder of the study. At the 4 hr. and 8 hr. timepoints, the animals were administered antitoxin as described above. On day 4, at the 0 hr. timepoint, each animal was again administered antitoxin as described above. At 1 hr., each animal was orally challenged with 1 ml of C. difficile inoculum, which contained approx. 100 C. difficile strain 43596 organisms in sterile saline. C. difficile strain 43596, which is a serogroup C strain, was chosen because it is representative of one of the most frequently-occurring serogroups isolated from patients with antibiotic-associated pseudomembranous colitis. [Delmee et al., J. Clin. Microbiol., 28:2210-2214 (1985).] In addition, this strain has been previously demonstrated to be virulent in the hamster model of infection. [Delmee and Avesani, J. Med. Microbiol., 33:85-90 (1990).] At the 4 hr. and 8 hr. timepoints, the animals were administered antitoxin as described above. On days 5 through 13, the animals were administered antitoxin 3x per day as described for day 1 above, and observed for the onset of diarrhea and death. On the morning of day 14, the final results of the study were tabulated. These results are shown in Table 13.

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Representative animals from those that died in the Pre-Immune and CTA groups were necropsied. Viable C. difficile organisms were cultured from the ceca of these animals, and the gross pathology of the gastrointestinal tracts of these animals was consistent with that expected for C. difficile disease (inflamed, distended, hemorrhagic cecum, filled with watery diarrhea-like material). In addition, the clindamycin control animal remained healthy throughout the entire study period,

therefore indicating that the hamsters used in the study had not previously been colonized with endogenous *C. difficile* organisms prior to the start of the study. Following the final antitoxin treatment on day 13, a single surviving animal from the CTA group, and also from the CTAB group, was sacrificed and necropsied. No pathology was noted in either animal.

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TABLE 13
Treatment Results

Treatment Group	No. Animals Surviving	No. Animals Dead
Pre-Immune	.1	6
CTA (Antitoxin A only)	5	2
CTAB (Antitoxin A + Antitoxin B)	7	0

Treatment of hamsters with orally-administered toxin A and toxin B antitoxin (group CTAB) successfully protected 7 out of 7 (100%) of the animals from C. difficile disease. Treatment of hamsters with orally-administered toxin A antitoxin (group CTA) protected 5 out of 7 (71%) of these animals from C. difficile disease. Treatment using pre-immune IgY was not protective against C. difficile disease, as only 1 out of 7 (14%) of these animals survived. These results demonstrate that the avian toxin A antitoxin and the avian toxin A + toxin B antitoxin effectively protected the hamsters from C. difficile disease. These results also suggest that although the neutralization of toxin A alone confers some degree of protection against C. difficile disease, in order to achieve maximal protection, simultaneous antitoxin A and antitoxin B activity is necessary.

c) Long-Term Survival Of Treated Hamsters

It has been previously reported in the literature that hamsters treated with orally-administered bovine antitoxin IgG concentrate are protected from *C. difficile* disease as long as the treatment is continued, but when the treatment is stopped, the animals develop diarrhea and subsequently die within 72 hrs. [Lyerly et al., Infect. Immun., 59(6):2215-2218 (1991).]

In order to determine whether treatment of *C. difficile* disease using avian antitoxins promotes long-term survival following the discontinuation of treatment, the 4 surviving animals in group CTA, and the 6 surviving animals in group CTAB were observed for a period of 11 days (264 hrs.) following the discontinuation of antitoxin treatment described in section (b) above. All hamsters remained healthy through the entire post-treatment period. This result demonstrates that not only does treatment with avian antitoxin protect against the onset of *C. difficile* disease (*i.e.*, it is effective as a prophylactic), it also promotes long-term survival beyond the treatment period, and thus provides a lasting cure.

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EXAMPLE 10

In vivo Treatment Of Established C. difficile Infection In Golden

Syrian Hamsters With Avian Antitoxins Against C. difficile Toxins A And B

The ability of the avian C. difficile antitoxins, described in Example 8 above, to treat an established C. difficile infection was evaluated using the Golden Syrian hamster model. The Example involved (a) preparation of the avian C. difficile antitoxins, (b) in vivo treatment of hamsters with established C. difficile infection, and (c) histologic evaluation of cecal tissue.

a) Preparation Of The Avian C. difficile Antitoxins

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Eggs were collected from hens in group CTAB described in Example 8 (b) above, which were immunized with *C. difficile* toxoids and native toxins A and B. Eggs purchased from a local supermarket were used as a pre-immune (negative) control. Egg yolk immunoglobulin (IgY) was extracted from the 2 groups of eggs as described in Example 1 (c), and the final IgY pellets were solubilized in one-fourth the original yolk volume of Ensure® nutritional formula.

b) In vivo Treatment Of Hamsters With Established C. difficile Infection

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The avian C. difficile antitoxins prepared in section (a) above were evaluated for the ability to treat established C. difficile infection in hamsters using an animal model system which was modified from the procedure which was described for the hamster protection study in Example 8(b) above.

For the study, four separate experimental groups were used, with each group consisting of 7 female Golden Syrian hamsters (Charles River), approx. 10 weeks old, weighing approximately 100 gms. each. Each animal was housed separately, and was offered food and water *ad libitum* through the entire length of the study.

On day 1 of the study, the animals in all four groups were each predisposed to C. difficile infection by the oral administration of 3.0 mg of clindamycin-HCl (Sigma) in 1 ml of water.

On day 2, each animal in all four groups was orally challenged with 1 ml of C. difficile inoculum, which contained approximately 100 C. difficile strain 43596 organisms in sterile saline. C. difficile strain 43596 was chosen because it is representative of one of the most frequently-occurring serogroups isolated from patients with antibiotic-associated pseudomembranous colitis. [Delmee et al., J. Clin. Microbiol., 28:2210-2214 (1990).] In addition, as this was the same C. difficile strain used in all of the previous Examples above, it was again used in order to provide experimental continuity.

On day 3 of the study (24 hrs. post-infection), treatment was started for two of the four groups of animals. Each animal of one group was orally administered 1.0 ml of the CTAB IgY preparation (prepared in section (a) above) at the following timepoints: 0 hrs., 4 hrs., and 8 hrs. The animals in this group were designated "CTAB-24." The animals in the second group were each orally administered 1.0 ml of the pre-immune IgY preparation (also prepared in section (a) above) at the same timepoints as for the CTAB group. These animals were designated "Pre-24." Nothing was done to the remaining two groups of animals on day 3.

On day 4, 48 hrs. post-infection, the treatment described for day 3 above was repeated for the CTAB-24 and Pre-24 groups, and was initiated for the remaining two groups at the same timepoints. The final two groups of animals were designated "CTAB-48" and "Pre-48" respectively.

On days 5 through 9, the animals in all four groups were administered antitoxin or pre-immune IgY, 3x per day, as described for day 4 above. The four experimental groups are summarized in Table 14.

TABLE 14
Experimental Treatment Groups

Group Designation	Experimental Treatment	
CTAB-24	Infected, treatment w/antitoxin IgY started @ 24 hrs. post-infection.	
Pre-24	Infected, treatment w/pre-immune IgY started @ 24 hrs. post-infection.	
CTAB-48	Infected, treatment w/antitoxin IgY started @ 48 hrs. post-infection.	
Pre-48	Infected, treatment w/pre-immune IgY started @ 48 hrs. post-infection.	

All animals were observed for the onset of diarrhea and death through the conclusion of the study on the morning of day 10. The results of this study are displayed in Table 15.

TABLE 15
Experimental Outcome--Day 10

Treatment Group	No. Animals Surviving	No. Animals Dead
CTAB-24	6	l
Pre-24	0	7
CTAB-48	4	3
Pre-48	2	Š

Eighty-six percent of the animals which began receiving treatment with antitoxin IgY at 24 hrs. post-infection (CTAB-24 above) survived, while 57% of the animals treated with antitoxin IgY starting 48 hrs. post-infection (CTAB-48 above) survived. In contrast, none of the animals receiving pre-immune IgY starting 24 hrs.

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post-infection (Pre-24 above) survived, and only 29% of the animals which began receiving treatment with pre-immune IgY at 48 hrs. post-infection (Pre-48 above) survived through the conclusion of the study. These results demonstrate that avian antitoxins raised against *C. difficile* toxins A and B are capable of successfully treating established *C. difficile* infections in vivo.

c) Histologic Evaluation Of Cecal Tissue

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In order to further evaluate the ability of the IgY preparations tested in this study to treat established *C. difficile* infection, histologic evaluations were performed on cecal tissue specimens obtained from representative animals from the study described in section (b) above.

Immediately following death, cecal tissue specimens were removed from animals which died in the Pre-24 and Pre-48 groups. Following the completion of the study, a representative surviving animal was sacrificed and cecal tissue specimens were removed from the CTAB-24 and CTAB-48 groups. A single untreated animal from the same shipment as those used in the study was also sacrificed and a cecal tissue specimen was removed as a normal control. All tissue specimens were fixed overnight at 4°C in 10% buffered formalin. The fixed tissues were paraffin-embedded, sectioned, and mounted on glass microscope slides. The tissue sections were then stained using hematoxylin and eosin (H and E stain), and were examined by light microscopy.

Upon examination, the tissues obtained from the CTAB-24 and CTAB-48 animals showed no pathology, and were indistinguishable from the normal control. This observation provides further evidence for the ability of avian antitoxins raised against *C. difficile* toxins A and B to effectively treat established *C. difficile* infection, and to prevent the pathologic consequences which normally occur as a result of *C. difficile* disease.

In contrast, characteristic substantial mucosal damage and destruction was observed in the tissues of the animals from the Pre-24 and Pre-48 groups which died

from C. difficile disease. Normal tissue architecture was obliterated in these two preparations, as most of the mucosal layer was observed to have sloughed away, and there were numerous large hemorrhagic areas containing massive numbers of erythrocytes.

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EXAMPLE 11

Cloning And Expression Of C. difficile Toxin A Fragments

The toxin A gene has been cloned and sequenced, and shown to encode a protein of predicted MW of 308 kd. [Dove et al., Infect. Immun., 58:480-488 (1990).] Given the expense and difficulty of isolating native toxin A protein, it would be advantageous to use simple and inexpensive procaryotic expression systems to produce and purify high levels of recombinant toxin A protein for immunization purposes. Ideally, the isolated recombinant protein would be soluble in order to preserve native antigenicity, since solubilized inclusion body proteins often do not fold into native conformations. To allow ease of purification, the recombinant protein should be expressed to levels greater than 1 mg/liter of E. coli culture.

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To determine whether high levels of recombinant toxin A protein can be produced in *E. coli*, fragments of the toxin A gene were cloned into various prokaryotic expression vectors, and assessed for the ability to express recombinant toxin A protein in *E. coli*. Three prokaryotic expression systems were utilized. These systems were chosen because they drive expression of either fusion (pMALc and pGEX2T) or native (pET23a-c) protein to high levels in *E. coli*, and allow affinity purification of the expressed protein on a ligand containing column. Fusion proteins expressed from pGEX vectors bind glutathione agarose beads, and are eluted with reduced glutathione. pMAL fusion proteins bind amylose resin, and are eluted with maltose. A poly-histidine tag is present at either the N-terminal (pET16b) or C-terminal (pET23a-c) end of pET fusion proteins. This sequence specifically binds Ni₂* chelate columns, and is eluted with imidazole salts. Extensive descriptions of these

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vectors are available [Williams et al. (1995) DNA Cloning 2: Expression Systems, Glover and Hames, eds. IRL Press, Oxford, pp. 15-58], and will not be discussed in detail here. The Example involved (a) cloning of the toxin A gene, (b) expression of large fragments of toxin A in various prokaryotic expression systems, (c) identification of smaller toxin A gene fragments that express efficiently in E. coli, (d) purification of recombinant toxin A protein by affinity chromatography, and (e) demonstration of functional activity of a recombinant fragment of the toxin A gene.

a) Cloning Of The Toxin A Gene

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A restriction map of the toxin A gene is shown in Figure 6. The encoded protein contains a carboxy terminal ligand binding region, containing multiple repeats of a carbohydrate binding domain. [von Eichel-Streiber and Sauerborn, Gene 96:107-113 (1990).] The toxin A gene was cloned in three pieces, by using either the polymerase chain reaction (PCR) to amplify specific regions, (regions 1 and 2, Figure 6) or by screening a constructed genomic library for a specific toxin A gene fragment (region 3, Figure 6). The sequences of the utilized PCR primers are P1: 5' GGAAATT TAGCTGCAGCATCTGAC 3' (SEQ ID NO.:1); P2: 5' TCTAGCAAATTCGCTTGT GTTGAA 3' (SEQ ID NO.:2); P3: 5' CTCGCATATAGCATTAGACC 3' (SEQ ID NO.:3); and P4: 5' CTATCTAGGCCTAAAGTAT 3' (SEQ ID NO.:4). These regions were cloned into prokaryotic expression vectors that express either fusion (pMALc and pGEX2T) or native (pET23a-c) protein to high levels in *E. coli*, and allow affinity purification of the expressed protein on a ligand containing column.

Clostridium difficile VPI strain 10463 was obtained from the ATCC (ATCC #43255) and grown under anaerobic conditions in brain-heart infusion medium (BBL). High molecular-weight C. difficile DNA was isolated essentially as described by Wren and Tabaqchali (1987) J. Clin. Microbiol., 25:2402, except proteinase K and sodium dodecyl sulfate (SDS) was used to disrupt the bacteria, and cetyltrimethylammonium bromide precipitation [as described in Ausubel et al., Current Protocols in Molecular Biology (1989)] was used to remove carbohydrates from the cleared lysate. The

integrity and yield of genomic DNA was assessed by comparison with a serial dilution of uncut lambda DNA after electrophoresis on an agarose gel.

Fragments 1 and 2 were cloned by PCR, utilizing a proofreading thermostable DNA polymerase (native pfu polymerase; Stratagene). The high fidelity of this polymerase reduces the mutation problems associated with amplification by error prone polymerases (e.g., Taq polymerase). PCR amplification was performed using the indicated PCR primers (Figure 6) in 50 µl reactions containing 10 mM Tris-HCl(8.3), 50 mM KCl, 1.5 mM MgCl₂, 200 μ M each dNTP, 0.2 μ M each primer, and 50 ng C. difficile genomic DNA. Reactions were overlaid with 100 µl mineral oil, heated to 94°C for 4 min, 0.5 µl native pfu polymerase (Stratagene) added, and the reaction cycled 30x at 94°C for 1 min, 50°C for 1 min, 72°C for 4 min, followed by 10 min at 72°C. Duplicate reactions were pooled, chloroform extracted, and ethanol precipitated. After washing in 70% ethanol, the pellets were resuspended in 50 µl TE buffer [10 mM Tris-HCL, 1 mM EDTA pH 8.0]. Aliquots of 10µl each were restriction digested with either EcoRI/HincII (fragment 1) or EcoRI/PstI (fragment 2), and the appropriate restriction fragments were gel purified using the Prep-A-Gene kit (BioRad), and ligated to either EcoRI/Smal-restricted pGEX2T (Pharmacia) vector (fragment 1), or the EcoRI/Pstl pMAlc (New England Biolabs) vector (fragment 2). Both clones are predicted to produce in-frame fusions with either the glutathione-S-transferase protein (pGEX vector) or the maltose binding protein (pMAL vector). Recombinant clones were isolated, and confirmed by restriction digestion, using standard recombinant molecular biology techniques. [Sambrook et al., Molecular Cloning, A Laboratory Manual (1989), and designated pGA30-660 and pMA660-1100, respectively (see Figure 6 for description of the clone designations).]

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Fragment 3 was cloned from a genomic library of size selected *Pst*I digested *C. difficile* genomic DNA, using standard molecular biology techniques (Sambrook *et al.*). Given that the fragment 3 internal *Pst*I site is protected from cleavage in *C. difficile* genomic DNA [Price *et al.*, Curr. Microbiol., 16:55-60 (1987)], a 4.7 kb fragment from *Pst*I restricted *C. difficile* genomic DNA was gel purified, and ligated to

PstI restricted, phosphatase treated pUC9 DNA. The resulting genomic library was screened with a oligonucleotide primer specific to fragment 3, and multiple independent clones were isolated. The presence of fragment 3 in several of these clones was confirmed by restriction digestion, and a clone of the indicated orientation (Figure 6) was restricted with BamHI/HindIII, the released fragment purified by gel electrophoresis, and ligated into similarly restricted pET23c expression vector DNA (Novagen). Recombinant clones were isolated, and confirmed by restriction digestion. This construct is predicted to create both a predicted in frame fusion with the pET protein leader sequence, as well as a predicted C-terminal poly-histidine affinity tag, and is designated pPA1100-2680 (see Figure 6 for the clone designation).

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b) Expression Of Large Fragments Of Toxin A In E. coli

Protein expression from the three expression constructs made in (a) was induced, and analyzed by Western blot analysis with an affinity purified, goat polyclonal antiserum directed against the toxin A toxoid (Tech Lab). The procedures utilized for protein induction, SDS-PAGE, and Western blot analysis are described in detail in Williams et al (1995), supra. In brief, 5 ml 2X YT (16 g tryptone, 10 g yeast extract, 5 g NaCl per liter, pH 7.5 + 100 μg/ml ampicillin were added to cultures of bacteria (BL21 for pMAl and pGEX plasmids, and BL21(DE3)LysS for pET plasmids) containing the appropriate recombinant clone which were induced to express recombinant protein by addition of IPTG to 1 mM. Cultures were grown at 37°C, and induced when the cell density reached 0.5 OD600. Induced protein was allowed to accumulate for two hrs after induction. Protein samples were prepared by pelleting 1 ml aliquots of bacteria by centrifugation (1 min in a microfuge), and resuspension of the pelleted bacteria in 150 µl of 2x SDS-PAGE sample buffer [Williams et al. (1995), supra]. The samples were heated to 95°C for 5 min, the cooled and 5 or 10 µl aliquots loaded on 7.5% SDS-PAGE gels. BioRad high molecular weight protein markers were also loaded, to allow estimation of the MW of identified fusion proteins. After electrophoresis, protein was detected either generally by staining gels with Coomassie

blue, or specifically, by blotting to nitrocellulose for Western blot detection of specific immunoreactive protein. Western blots, (performed as described in Example 3) which detect toxin A reactive protein in cell lysates of induced protein from the three expression constructs are shown in Figure 7. In this figure, lanes 1-3 contain cell lysates prepared from *E. coli* strains containing pPA1100-2860 in B121(DE3)lysE cells; lanes 4-6 contain cell lysates prepared from *E. coli* strains containing pPA1100-2860 in B121(DE3)lysS cells; lanes 7-9 contain cell lysates prepared from *E. coli* strains containing pMA30-660; lanes 10-12 contain cell lysates prepared from *E. coli* strains containing pMA660-1100. The lanes were probed with an affinity purified goat antitoxin A polyclonal antibody (Tech Lab). Control lysates from uninduced cells (lanes 1, 7, and 10) contain very little immunoreactive material compared to the induced samples in the remaining lanes. The highest molecular weight band observed for each clone is consistent with the predicted size of the full length fusion protein.

Each construct directs expression of high molecular weight (HMW) protein that is reactive with the toxin A antibody. The size of the largest immunoreactive bands from each sample is consistent with predictions of the estimated MW of the intact fusion proteins. This demonstrates that the three fusions are in-frame, and that none of the clones contain cloning artifacts that disrupt the integrity of the encoded fusion protein. However, the Western blot demonstrates that fusion protein from the two larger constructs (pGA30-660 and pPA1100-2680) are highly degraded. Also, expression levels of toxin A proteins from these two constructs are low, since induced protein bands are not visible by Coomassie staining (not shown). Several other expression constructs that fuse large sub-regions of the toxin A gene to either pMALc or pET23a-c expression vectors, were constructed and tested for protein induction. These constructs were made by mixing gel purified restriction fragments, derived from the expression constructs shown in Figure 6, with appropriately cleaved expression vectors, ligating, and selecting recombinant clones in which the toxin A restriction fragments had ligated together and into the expression vector as predicted for in-frame fusions. The expressed toxin A interval within these constructs are shown in Figure 8, as well as the internal restriction sites utilized to make these constructs.

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As used herein, the term "interval" refers to any portion (i.e., any segment of the toxin which is less than the whole toxin molecule) of a clostridial toxin. In a preferred embodiment, "interval" refers to portions of C. difficile toxins such as toxin A or toxin B. It is also contemplated that these intervals will correspond to epitopes of immunologic importance, such as antigens or immunogens against which a neutralizing antibody response is effected. It is not intended that the present invention be limited to the particular intervals or sequences described in these Examples. It is also contemplated that sub-portions of intervals (e.g., an epitope contained within one interval or which bridges multiple intervals) be used as compositions and in the methods of the present invention.

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In all cases, Western blot analysis of each of these constructs with goat antitoxin A antibody (Tech Lab) detected HMW fusion protein of the predicted size (not shown). This confirms that the reading frame of each of these clones is not prematurely terminated, and is fused in the correct frame with the fusion partner. However, the Western blot analysis revealed that in all cases, the induced protein is highly degraded, and, as assessed by the absence of identifiable induced protein bands by Coomassie Blue staining, are expressed only at low levels. These results suggest that expression of high levels of intact toxin A recombinant protein is not possible when large regions of the toxin A gene are expressed in *E. coli* using these expression vectors.

c) High Level Expression Of Small Toxin A Protein Fusions In E. coli

Experience indicates that expression difficulties are often encountered when large (greater than 100 kd) fragments are expressed in *E. coli*. A number of expression constructs containing smaller fragments of the toxin A gene were constructed, to determine if small regions of the gene can be expressed to high levels without extensive protein degradation. A summary of these expression constructs are shown in Figure 9. All were constructed by in-frame fusions of convenient toxin A

restriction fragments to either the pMALc or pET23a-c vectors. Protein preparations from induced cultures of each of these constructs were analyzed by both Coomassie Blue staining and Western analysis as in (b) above. In all cases, higher levels of intact, full length fusion proteins were observed than with the larger recombinants from section (b).

d) Purification Of Recombinant Toxin A Protein

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Large scale (500 ml) cultures of each recombinant from (c) were grown, induced, and soluble and insoluble protein fractions were isolated. The soluble protein extracts were affinity chromatographed to isolate recombinant fusion protein, as described [Williams et al. (1994), supra]. In brief, extracts containing tagged pET fusions were chromatographed on a nickel chelate column, and eluted using imidazole salts as described by the distributor (Novagen). Extracts containing soluble pMAL fusion protein were prepared and chromatographed in column buffer (10 mM NaPO₄, 0.5M NaCl, 10 mM β-mercaptoethanol, pH 7.2) over an amylose resin column (New England Biolabs), and eluted with column buffer containing 10 mM maltose as described [Williams et al. (1995), supra]. When the expressed protein was found to be predominantly insoluble, insoluble protein extracts were prepared by the method described in Example 17, infra. The results are summarized in Table 16. Figure 10 shows the sample purifications of recombinant toxin A protein. In this figure, lanes 1 and 2 contain MBP fusion protein purified by affinity purification of soluble protein.

TABLE 16
Purification Of Recombinant Toxin A Protein

friends and the	The state of the s				
Clone (4)	Protein Solubility	Yield Affinity Purified Soluble Protein ⁽⁹⁾	% Intact Soluble Fusion Protein ^(c)	Yield Intact Insoluble Fusion Protein	
pMA30-270	Soluble	4 mg/500 mls	10%	NA	
PMA30-300	Soluble	4 mg/500 mls	5–10%	NA	
pMA300-660	Insoluble		NA	10 mg/500 ml	
pMA660-1100	Soluble	4.5 mg/500 mls	50%	NA	
pMA1100-1610	Soluble	18 mg/500 mls	10%	NA	
pMA1610-1870	Both	22 mg/500 mls	90%	20 mg/500 ml	
pMA1450-1870	Insoluble		NA	0.2 mg/500 ml	
pPA1100-1450	Soluble	0.1 mg/500 mls	90%	NA	
pPA1100-1870	Soluble	0.02 mg/500 mls	90%	NA	
pMA1870-2680	Both	12 mg/500 mls	80%	NA	
pPa1870-2680	Insoluble		NA	10 mg/500 ml	

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pP = pET23 vector, pM=pMALc vector, A=toxin A.

Based on 1.5 OD₂₈₀ = 1 mg/ml (extinction coefficient of MBP).

(c) Estimated by Coomassie staining of SDS-PAGE gels.

Lanes 3 and 4 contain MBP fusion protein purified by solubilization of insoluble inclusion bodies. The purified fusion protein samples are pMA1870-2680 (lane 1), pMA660-1100 (lane 2), pMA300-600 (lane 3) and pMA1450-1870 (lane 4).

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Poor yields of affinity purified protein were obtained when poly-histidine tagged pET vectors were used to drive expression (pPA1100-1450, pP1100-1870). However, significant protein yields were obtained from pMAL expression constructs spanning the entire toxin A gene, and yields of full-length soluble fusion protein ranged from an estimated 200-400 µg/500 ml culture (pMA30-300) to greater than 20 mg/500 ml culture (pMA1610-1870). Only one interval was expressed to high levels as strictly insoluble protein (pMA300-660). Thus, although high level expression was not observed when using large expression constructs from the toxin A gene, usable levels of recombinant protein spanning the entire toxin A gene were obtainable by

isolating induced protein from a series of smaller pMAL expression constructs that span the entire toxin A gene. This is the first demonstration of the feasibility of expressing recombinant toxin A protein to high levels in E. coli.

e) Hemagglutination Assay Using The Toxin A Recombinant Proteins

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The carboxy terminal end consisting of the repeating units contains the hemagglutination activity or binding domain of C. difficile toxin A. To determine whether the expressed toxin A recombinants retain functional activity, hemagglutination assays were performed. Two toxin A recombinant proteins, one containing the binding domain as either soluble affinity purified protein (pMA1870-2680) or SDS solubilized inclusion body protein (pPA1870-2680) and soluble protein from one region outside that domain (pMA1100-1610) were tested using a described procedure. [H.C. Krivan et. al., Infect. Immun., 53:573 (1986).] Citrated rabbit red blood cells (RRBC)(Cocalico) were washed several times with Tris-buffer (0.1M Tris and 50 mM NaCl) by centrifugation at 450 x g for 10 minutes at 4° C. A 1% RRBC suspension was made from the packed cells and resuspended in Tris-buffer. Dilutions of the recombinant proteins and native toxin A (Tech Labs) were made in the Trisbuffer and added in duplicate to a round-bottomed 96-well microtiter plate in a final volume of 100 µl. To each well, 50 µl of the 1% RRBC suspension was added, mixed by gentle tapping, and incubated at 4°C for 3-4 hours. Significant hemagglutination occurred only in the recombinant proteins containing the binding domain (pMA 1870-2680) and native toxin A. The recombinant protein outside the binding domain (pMA 1100-1610) displayed no hemagglutination activity. Using equivalent protein concentrations, the hemagglutination titer for toxin A was 1:256, while titers for the soluble and insoluble recombinant proteins of the binding domain were 1:256 and about 1:5000. Clearly, the recombinant proteins tested retained functional activity and were able to bind RRBC's.

EXAMPLE 12

Functional Activity Of IgY Reactive Against Toxin A Recombinants

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The expression of recombinant toxin A protein as multiple fragments in E. coli has demonstrated the feasibility of generating toxin A antigen through use of recombinant methodologies (Example 11). The isolation of these recombinant proteins allows the immunoreactivity of each individual subregion of the toxin A protein to be determined (i.e., in a antibody pool directed against the native toxin A protein). This identifies the regions (if any) for which little or no antibody response is elicited when the whole protein is used as a immunogen. Antibodies directed against specific fragments of the toxin A protein can be purified by affinity chromatography against recombinant toxin A protein, and tested for neutralization ability. This identifies any toxin A subregions that are essential for producing neutralizing antibodies. Comparison with the levels of immune response directed against these intervals when native toxin is used as an immunogen predicts whether potentially higher titers of neutralizing antibodies can be produced by using recombinant protein directed against a individual region, rather than the entire protein. Finally, since it is unknown whether antibodies reactive to the recombinant toxin A proteins produced in Example 11 neutralize toxin A as effectively as antibodies raised against native toxin A (Examples 9 and 10), the protective ability of a pool of antibodies affinity purified against recombinant toxin A fragments was assessed for its ability to neutralize toxin A.

This Example involved (a) epitope mapping of the toxin A protein to determine the titre of specific antibodies directed against individual subregions of the toxin A protein when native toxin A protein is used as an immunogen, (b) affinity purification of IgY reactive against recombinant proteins spanning the toxin A gene, (c) toxin A neutralization assays with affinity purified IgY reactive to recombinant toxin A protein to identify subregions of the toxin A protein that induce the production of neutralizing antibodies, and determination of whether complete neutralization of toxin A can be elicited with a mixture of antibodies reactive to recombinant toxin A protein.

a) Epitope Mapping Of The Toxin A Gene

The affinity purification of recombinant toxin A protein specific to defined intervals of the toxin A protein allows epitope mapping of antibody pools directed against native toxin A. This has not previously been possible, since previous expression of toxin A recombinants has been assessed only by Western blot analysis, without knowledge of the expression levels of the protein [e.g., von Eichel-Streiber et al, J. Gen. Microbiol., 135:55-64 (1989)]. Thus, high or low reactivity of recombinant toxin A protein on Western blots may reflect protein expression level differences, not immunoreactivity differences. Given that the purified recombinant protein generated in Example 11 have been quantitated, the issue of relative immunoreactivity of individual regions of the toxin A protein was precisely addressed.

For the purposes of this Example, the toxin A protein was subdivided into 6 intervals (1-6), numbered from the amino (interval 1) to the carboxyl (interval 6) termini.

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The recombinant proteins corresponding to these intervals were from expression clones (see Example 11(d) for clone designations) pMA30-300 (interval 1), pMA300-660 (interval 2), pMA660-1100 (interval 3), pPA1100-1450 (interval 4), pMA1450-1870 (interval 5) and pMA1870-2680 (interval 6). These 6 clones were selected because they span the entire protein from amino acids numbered 30 through 2680, and subdivide the protein into 6 small intervals. Also, the carbohydrate binding repeat interval is contained specifically in one interval (interval 6), allowing evaluation of the immune response specifically directed against this region. Western blots of 7.5% SDS-PAGE gels, loaded and electrophoresed with defined quantities of each recombinant protein, were probed with either goat antitoxin A polyclonal antibody (Tech Lab) or chicken antitoxin A polyclonal antibody [pCTA IgY, Example 8(c)]. The blots were prepared and developed with alkaline phosphatase as previously described [Williams et al. (1995), supra]. At least 90% of all reactivity, in either goat or chicken antibody pools, was found to be directed against the ligand binding domain (interval 6). The remaining immunoreactivity was directed against all five remaining

intervals, and was similar in both antibody pools, except that the chicken antibody showed a much lower reactivity against interval 2 than the goat antibody.

This clearly demonstrates that when native toxin A is used as an immunogen in goats or chickens, the bulk of the immune response is directed against the ligand binding domain of the protein, with the remaining response distributed throughout the remaining 2/3 of the protein.

b) Affinity Purification Of IgY Reactive Against Recombinant Toxin A Protein

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Affinity columns, containing recombinant toxin A protein from the 6 defined intervals in (a) above, were made and used to (i) affinity purify antibodies reactive to each individual interval from the CTA IgY preparation [Example 8(c)], and (ii) deplete interval specific antibodies from the CTA IgY preparation. Affinity columns were made by coupling 1 ml of PBS-washed Actigel resin (Sterogene) with region specific protein and 1/10 final volume of Ald-coupling solution (1M sodium cyanoborohydride). The total region specific protein added to each reaction mixture was 2.7 mg (interval 1), 3 mg (intervals 2 and 3), 0.1 mg (interval 4), 0.2 mg (interval 5) and 4 mg (interval 6). Protein for intervals 1, 3, and 6 was affinity purified pMAI fusion protein in column buffer (see Example 11). Interval 4 was affinity purified poly-histidine containing pET fusion in PBS; intervals 2 and 5 were from inclusion body preparations of insoluble pMAL fusion protein, dialyzed extensively in PBS. Aliquots of the supernatants from the coupling reactions, before and after coupling. were assessed by Coomassie staining of 7.5% SDS-PAGE gels. Based on protein band intensities, in all cases greater than 50% coupling efficiencies were estimated. The resins were poured into 5 ml BioRad columns, washed extensively with PBS, and stored at 4°C.

Aliquots of the CTA IgY polyclonal antibody preparation were depleted for each individual region as described below. A 20 ml sample of the CTA IgY preparation [Example 8(c)] was dialyzed extensively against 3 changes of PBS (1 liter for each dialysis), quantitated by absorbance at OD₂₈₀, and stored at 4°C. Six 1 ml

aliquots of the dialyzed IgY preparation were removed, and depleted individually for each of the six intervals. Each 1 ml aliquot was passed over the appropriate affinity column, and the eluate twice reapplied to the column. The eluate was collected, and pooled with a 1 ml PBS wash. Bound antibody was eluted from the column by washing with 5 column volumes of 4 M Guanidine-HCl (in 10 mM Tris-HCl, pH 8.0). The column was reequilibrated in PBS, and the depleted antibody stock reapplied as described above. The eluate was collected, pooled with a 1 ml PBS wash, quantitated by absorbance at OD₂₈₀, and stored at 4° C. In this manner, 6 aliquots of the CTA IgY preparation were individually depleted for each of the 6 toxin A intervals, by two rounds of affinity depletion. The specificity of each depleted stock was tested by Western blot analysis. Multiple 7.5% SDS-PAGE gels were loaded with protein samples corresponding to all 6 toxin A subregions. After electrophoresis, the gels were blotted, and protein transfer confirmed by Ponceau S staining [protocols described in Williams et al. (1995), supra]. After blocking the blots 1 hr at 20°C in PBS+ 0.1% Tween 20 (PBST) containing 5% milk (as a blocking buffer), 4 ml of either a 1/500 dilution of the dialyzed CTA IgY preparation in blocking buffer, or an equivalent amount of the six depleted antibody stocks (using OD₂₈₀ to standardize antibody concentration) were added and the blots incubated a further 1 hr at room temperature. The blots were washed and developed with alkaline phosphatase (using a rabbit anti-chicken alkaline phosphate conjugate as a secondary antibody) as previously described [Williams et al. (1995), supra]. In all cases, only the target interval was depleted for antibody reactivity, and at least 90% of the reactivity to the target intervals was specifically depleted.

Region specific antibody pools were isolated by affinity chromatography as described below. Ten mls of the dialyzed CTA IgY preparation were applied sequentially to each affinity column, such that a single 10 ml aliquot was used to isolate region specific antibodies specific to each of the six subregions. The columns were sequentially washed with 10 volumes of PBS, 6 volumes of BBS-Tween, 10 volumes of TBS, and eluted with 4 ml Actisep elution media (Sterogene). The eluate was dialyzed extensively against several changes of PBS, and the affinity purified

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antibody collected and stored at 4°C. The volumes of the eluate increased to greater than 10 mls during dialysis in each case, due to the high viscosity of the Actisep elution media. Aliquots of each sample were 20x concentrated using Centricon 30 microconcentrators (Amicon) and stored at 4°C. The specificity of each region specific antibody pool was tested, relative to the dialyzed CTA IgY preparation, by Western blot analysis, exactly as described above, except that 4 ml samples of blocking buffer containing 100 µl region specific antibody (unconcentrated) were used instead of the depleted CTA IgY preparations. Each affinity purified antibody preparation was specific to the defined interval, except that samples purified against intervals 1-5 also reacted with interval 6. This may be due to non-specific binding to the interval 6 protein, since this protein contains the repetitive ligand binding domain which has been shown to bind antibodies nonspecifically. [Lyerly et al., Curr. Microbiol., 19:303-306 (1989).]

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The reactivity of each affinity purified antibody preparation to the corresponding proteins was approximately the same as the reactivity of the 1/500 diluted dialyzed CTA IgY preparation standard. Given that the specific antibody stocks were diluted 1/40, this would indicate that the unconcentrated affinity purified antibody stocks contain 1/10-1/20 the concentration of specific antibodies relative to the starting CTA IgY preparation.

c) Toxin A Neutralization Assay Using Antibodies Reactive Toward Recombinant Toxin A Protein

The CHO toxin neutralization assay [Example 8(d)] was used to assess the ability of the depleted or enriched samples generated in (b) above to neutralize the cytotoxicity of toxin A. The general ability of affinity purified antibodies to neutralize toxin A was assessed by mixing together aliquots of all 6 concentrated stocks of the 6 affinity purified samples generated in (b) above, and testing the ability of this mixture to neutralize a toxin A concentration of 0.1 µg/ml. The results, shown in Figure 11, demonstrate almost complete neutralization of toxin A using the affinity purified (AP) mix. Some epitopes within the recombinant proteins utilized for affinity purification

were probably lost when the proteins were denatured before affinity purification [by Guanidine-HCl treatment in (b) above]. Thus, the neutralization ability of antibodies directed against recombinant protein is probably underestimated using these affinity purified antibody pools. This experiment demonstrates that antibodies reactive to recombinant toxin A can neutralize cytotoxicity, suggesting that neutralizing antibodies may be generated by using recombinant toxin A protein as immunogen.

In view of the observation that the recombinant expression clones of the toxin A gene divide the protein into 6 subregions, the neutralizing ability of antibodies directed against each individual region was assessed. The neutralizing ability of antibodies directed against the ligand binding domain of toxin A was determined first.

In the toxin neutralization experiment shown in Figure 11, interval 6 specific antibodies (interval 6 contains the ligand binding domain) were depleted from the dialyzed PEG preparation, and the effect on toxin neutralization assayed. Interval 6 antibodies were depleted either by utilizing the interval 6 depleted CTA IgY preparation from (b) above ("-6 aff. depleted" in Figure 11), or by addition of interval 6 protein to the CTA IgY preparation (estimated to be a 10 fold molar excess over anti-interval 6 immunoglobulin present in this preparation) to competitively compete for interval 6 protein ("-6 prot depleted" in Figure 11). In both instances, removal of interval 6 specific antibodies reduces the neutralization efficiency relative to the starting CTA IgY preparation. This demonstrates that antibodies directed against interval 6 contribute to toxin neutralization. Since interval 6 corresponds to the ligand binding domain of the protein, these results demonstrate that antibodies directed against this region in the PEG preparation contribute to the neutralization of toxin A in this assay. However, it is significant that after removal of these antibodies, the PEG preparation retains significant ability to neutralize toxin A (Figure 11). This neutralization is probably due to the action of antibodies specific to other regions of the toxin A protein, since at least 90% of the ligand binding region reactive antibodies were removed in the depleted sample prepared in (b) above. This conclusion was supported by comparison of the toxin neutralization of the affinity purified (AP) mix compared to affinity purified interval 6 antibody alone. Although some neutralization

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ability was observed with AP interval 6 antibodies alone, the neutralization was significantly less than that observed with the mixture of all 6 AP antibody stocks (not shown).

Given that the mix of all six affinity purified samples almost completely neutralized the cytotoxicity of toxin A (Figure 11), the relative importance of antibodies directed against toxin A intervals 1-5 within the mixture was determined. This was assessed in two ways. First, samples containing affinity purified antibodies representing 5 of the 6 intervals were prepared, such that each individual region was depleted from one sample. Figure 12 demonstrates a sample neutralization curve, comparing the neutralization ability of affinity purified antibody mixes without interval 4 (-4) or 5 (-5) specific antibodies, relative to the mix of all 6 affinity purified antibody stocks (positive control). While the removal of interval 5 specific antibodies had no effect on toxin neutralization (or intervals 1-3, not shown), the loss of interval 4 specific antibodies significantly reduced toxin neutralization (Figure 12).

Similar results were seen in a second experiment, in which affinity purified antibodies, directed against a single region, were added to interval 6 specific antibodies, and the effects on toxin neutralization assessed. Only interval 4 specific antibodies significantly enhanced neutralization when added to interval 6 specific antibodies (Figure 13). These results demonstrate that antibodies directed against interval 4 (corresponding to clone pPA1100-1450 in Figure 9) are important for neutralization of cytotoxicity in this assay. Epitope mapping has shown that only low levels of antibodies reactive to this region are generated when native toxin A is used as an immunogen [Example 12(a)]. It is hypothesized that immunization with recombinant protein specific to this interval will elicit higher titers of neutralizing antibodies. In summary, this analysis has identified two critical regions of the toxin A protein against which neutralizing antibodies are produced, as assayed by the CHO neutralization assay.

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EXAMPLE 13

Production And Evaluation Of Avian Antitoxin

Against C. difficile Recombinant Toxin A Polypeptide

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In Example 12, we demonstrated neutralization of toxin A mediated cytotoxicity by affinity purified antibodies reactive to recombinant toxin A protein. To determine whether antibodies <u>raised</u> against a recombinant polypeptide fragment of *C. difficile* toxin A may be effective in treating clostridial diseases, antibodies to recombinant toxin A protein representing the binding domain were generated. Two toxin A binding domain recombinant polypeptides, expressing the binding domain in either the pMALc (pMA1870-2680) or pET 23(pPA1870-2680) vector, were used as immunogens. The pMAL protein was affinity purified as a soluble product [Example 12(d)] and the pET protein was isolated as insoluble inclusion bodies [Example 12(d)] and solubilized to an immunologically active protein using a proprietary method described in a pending patent application (U.S. Patent Application Serial No. 08/129,027). This Example involves (a) immunization, (b) antitoxin collection, (c) determination of antitoxin antibody titer, (d) anti-recombinant toxin A neutralization of toxin A hemagglutination activity *in vitro*, and (e) assay of *in vitro* toxin A neutralizing activity.

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a) Immunization

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The soluble and the inclusion body preparations each were used separately to immunize hens. Both purified toxin A polypeptides were diluted in PBS and emulsified with approximately equal volumes of CFA for the initial immunization or IFA for subsequent booster immunizations. On day zero, for each of the recombinant preparations, two egg laying white Leghorn hens (obtained from local breeder) were each injected at multiple sites (intramuscular and subcutaneous) with 1 ml of recombinant adjuvant mixture containing approximately 0.5 to 1.5 mgs of recombinant toxin A. Booster immunizations of 1.0 mg were given on days 14 and day 28.

b) Antitoxin Collection

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Total yolk immune IgY was extracted as described in the standard PEG protocol (as in Example 1) and the final IgY pellet was dissolved in sterile PBS at the original yolk volume. This material is designated "immune recombinant IgY" or "immune IgY."

c) Antitoxin Antibody Titer

To determine if the recombinant toxin A protein was sufficiently immunogenic to raise antibodies in hens, the antibody titer of a recombinant toxin A polypeptide was determined by ELISA. Eggs from both hens were collected on day 32, the yolks pooled and the antibody was isolated using PEG as described. The immune recombinant IgY antibody titer was determined for the soluble recombinant protein containing the maltose binding protein fusion generated in p-Mal (pMA1870-2680). Ninety-six well Falcon Pro-bind plates were coated overnight at 4°C with 100 µl /well of toxin A recombinant at 2.5 µg /µl in PBS containing 0.05% thimerosal. Another plate was also coated with maltose binding protein (MBP) at the same concentration, to permit comparison of antibody reactivity to the fusion partner. The next day, the wells were blocked with PBS containing 1% bovine serum albumin (BSA) for 1 hour at 37°C. IgY isolated from immune or preimmune eggs was diluted in antibody diluent (PBS containing 1% BSA and 0.05% Tween-20), and added to the blocked wells and incubated for 1 hour at 37°C. The plates were washed three times with PBS with 0.05% Tween-20, then three times with PBS. Alkaline phosphatase conjugated rabbit anti-chicken IgG (Sigma) diluted 1:1000 in antibody diluent was added to the plate, and incubated for I hour at 37°C. The plates were washed as before and substrate was added, [p-nitrophenyl phosphate (Sigma)] at 1 mg/ml in 0.05M Na₂CO₃, pH 9.5 and 10 mM MgCl₂. The plates were evaluated quantitatively on a Dynatech MR 300 Micro EPA plate reader at 410 nm about 10 minutes after the addition of substrate.

Based on these ELISA results, high antibody titers were raised in chickens immunized with the toxin A recombinant polypeptide. The recombinant appeared to

be highly immunogenic, as it was able to generate high antibody titers relatively quickly with few immunizations. Immune IgY titer directed specifically to the toxin A portion of the recombinant was higher than the immune IgY titer to its fusion partner, the maltose binding protein, and significantly higher than the preimmune IgY. ELISA titers (reciprocal of the highest dilution of IgY generating a signal) in the preimmune IgY to the MBP or the recombinant was <1:30 while the immune IgY titers to MBP and the toxin A recombinant were 1:18750 and > 1:93750 respectively. Importantly, the anti-recombinant antibody titers generated in the hens against the recombinant polypeptide is much higher, compared to antibodies to that region raised using native toxin A. The recombinant antibody titer to region 1870-2680 in the CTA antibody preparation is at least five-fold lower compared to the recombinant generated antibodies (1:18750 versus >1:93750). Thus, it appears a better immune response can be generated against a specific recombinant using that recombinant as the immunogen compared to the native toxin A.

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This observation is significant, as it shows that because recombinant portions stimulate the production of antibodies, it is not necessary to use native toxin molecules to produce antitoxin preparations. Thus, the problems associated with the toxicity of the native toxin are avoided and large-scale antitoxin production is facilitated.

d) Anti-Recombinant Toxin A Neutralization Of Toxin A Hemagglutination Activity In Vitro

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Toxin A has hemagglutinating activity besides cytotoxic and enterotoxin properties. Specifically, toxin A agglutinates rabbit erythrocytes by binding to a trisaccharide (gal 1-3B1-4GlcNAc) on the cell surface. [H. Krivan et al., Infect. Immun., 53:573-581 (1986).] We examined whether the anti-recombinant toxin A (immune IgY, antibodies raised against the insoluble product expressed in pET) can neutralize the hemagglutination activity of toxin A in vitro. The hemagglutination assay procedure used was described by H.C. Krivan et al. Polyethylene glycol-fractionated immune or preimmune IgY were pre-absorbed with citrated rabbit erythrocytes prior to performing the hemagglutination assay because we have found

that IgY alone can agglutinate red blood cells. Citrated rabbit red blood cells (RRBC's)(Cocalico) were washed twice by centrifugation at 450 x g with isotonic buffer (0.1 M Tris-HCl, 0.05 M NaCl, pH 7.2). RRBC-reactive antibodies in the IgY were removed by preparing a 10% RRBC suspension (made by adding packed cells to immune or preimmune IgY) and incubating the mixture for 1 hour at 37°C. The RRBCs were then removed by centrifugation. Neutralization of the hemagglutination activity of toxin A by antibody was tested in round-bottomed 96-well microtiter plates. Twenty-five µl of toxin A (36 µg /ml) (Tech Lab) in isotonic buffer was mixed with an equal volume of different dilutions of immune or preimmune IgY in isotonic buffer, and incubated for 15 minutes at room temperature. Then, 50 μ l of a 1% RRBC suspension in isotonic buffer was added and the mixture was incubated for 3 hours at 4°C. Positive control wells containing the final concentration of 9 μg/ml of toxin A after dilution without IgY were also included. Hemagglutination activity was assessed visually, with a diffuse matrix of RRBC's coating the bottom of the well representing a positive hemagglutination reaction and a tight button of RRBC's at the bottom of the well representing a negative reaction. The anti-recombinant immune IgY neutralized toxin A hemagglutination activity, giving a neutralization titer of 1:8. However, preimmune IgY was unable to neutralize the hemagglutination ability of toxin A.

e) Assay Of In Vitro Toxin A Neutralizing Activity

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The ability of the anti-recombinant toxin A IgY (immune IgY antibodies raised against pMA1870-2680, the soluble recombinant binding domain protein expressed in pMAL, designated as Anti-tox. A-2 in Figure 14, and referred to as recombinant region 6) and pre-immune IgY, prepared as described in Example 8(c) above, to neutralize the cytotoxic activity of toxin A was assessed *in vitro* using the CHO cell cytotoxicity assay, and toxin A (Tech Lab) at a concentration of 0.1µg/ml, as described in Example 8(d) above. As additional controls, the anti-native toxin A IgY (CTA) and pre-immune IgY preparations described in Example 8(c) above were also tested. The results are shown in Figure 14.

The anti-recombinant toxin A IgY demonstrated only partial neutralization of the cytotoxic activity of toxin A, while the pre-immune IgY did not demonstrate any significant neutralizing activity.

EXAMPLE 14

In vivo Neutralization Of C. difficile Toxin A

The ability of avian antibodies (IgY) raised against recombinant toxin A binding domain to neutralize the enterotoxin activity of C. difficile toxin A was evaluated in vivo using Golden Syrian hamsters. The Example involved:

- (a) preparation of the avian anti-recombinant toxin A IgY for oral administration;
- (b) in vivo protection of harmsters from C. difficile toxin A enterotoxicity by treatment of toxin A with avian anti-recombinant toxin A IgY; and (c) histologic evaluation of hamster ceca.

a) Preparation Of The Avian Anti-Recombinant Toxin A IgY For Oral Administration

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Eggs were collected from hens which had been immunized with the recombinant C. difficile toxin A fragment pMA1870-2680 (described in Example 13, above). A second group of eggs purchased at a local supermarket was used as a pre-immune (negative) control. Egg yolk immunoglobulin (IgY) was extracted by PEG from the two groups of eggs as described in Example 8(c), and the final IgY pellets were solubilized in one-fourth the original yolk volume using 0.1M carbonate buffer (mixture of NaHCO₃ and Na₂CO₃), pH 9.5. The basic carbonate buffer was used in order to protect the toxin A from the acidic pH of the stomach environment.

b) In vivo Protection Of Hamsters Against C. difficile Toxin A Enterotoxicity By Treatment Of Toxin A With Avian Anti-recombinant Toxin A IgY

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In order to assess the ability of the avian anti-recombinant toxin A IgY, prepared in section (a) above to neutralize the *in vivo* enterotoxin activity of toxin A, an *in vivo* toxin neutralization model was developed using Golden Syrian hamsters. This model was based on published values for the minimum amount of toxin A required to elicit diarrhea (0.08 mg toxin A/Kg body wt.) and death (0.16 mg toxin A/Kg body wt.) in hamsters when administered orally (Lyerly *et al.* Infect. Immun., 47:349-352 (1985).

For the study, four separate experimental groups were used, with each group consisting of 7 female Golden Syrian hamsters (Charles River), approx. three and one-half weeks old, weighing approx. 50 gms each. The animals were housed as groups of 3 and 4, and were offered food and water ad libitum through the entire length of the study.

For each animal, a mixture containing either 10µg of toxin A (0.2 mg/Kg) or 30µg of toxin A (0.6 mg/Kg) (C. difficile toxin A was obtained from Tech Lab and 1 ml of either the anti-recombinant toxin A IgY or pre-immune IgY (from section (a) above) was prepared. These mixtures were incubated at 37°C for 60 min. and were then administered to the animals by the oral route. The animals were then observed for the onset of diarrhea and death for a period of 24 hrs. following the administration of the toxin A+IgY mixtures, at the end of which time, the following results were tabulated and shown in Table 17:

TABLE 17
Study Outcome At 24 Hours

Experimental Group	Study Outcome at 24 Hours		
experimental Group	Healthy ¹	Diamhea ²	Dead'
10 µg Toxin A + Antitoxin Against Interval 6	7	0	0
30 µg Toxin A + Antitoxin Against Interval 6	7	0	0
10 μg Toxin A + Pre-Immune Serum	0.	5	2
30 μg Toxin A + Pre-Immune	0	5	2

- Animals remained healthy through the entire 24 hour study period.
- Animals developed diarrhea, but did not die.

Animals developed diarrhea, and subsequently died.

Pretreatment of toxin A at both doses tested, using the anti-recombinant toxin A IgY, prevented all overt symptoms of disease in hamsters. Therefore, pretreatment of C. difficile toxin A, using the anti-recombinant toxin A IgY, neutralized the in vivo enterotoxin activity of the toxin A. In contrast, all animals from the two groups which received toxin A which had been pretreated using pre-immune IgY developed disease symptoms which ranged from diarrhea to death. The diarrhea which developed in the 5 animals which did not die in each of the two pre-immune groups, spontaneously resolved by the end of the 24 hr. study period.

c) Histologic Evaluation Of Hamster Ceca

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In order to further assess the ability of anti-recombinant toxin A IgY to protect hamsters from the enterotoxin activity of toxin A, histologic evaluations were performed on the ceca of hamsters from the study described in section (b) above.

Three groups of animals were sacrificed in order to prepare histological specimens. The first group consisted of a single representative animal taken from each of the 4 groups of surviving hamsters at the conclusion of the study described in section (b) above. These animals represented the 24 hr. timepoint of the study.

The second group consisted of two animals which were not part of the study described above, but were separately treated with the same toxin A + pre-immune IgY

mixtures as described for the animals in section (b) above. Both of these hamsters developed diarrhea, and were sacrificed 8 hrs. after the time of administration of the toxin A + pre-immune IgY mixtures. At the time of sacrifice, both animals were presenting symptoms of diarrhea. These animals represented the acute phase of the study.

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The final group consisted of a single untreated hamster from the same shipment of animals as those used for the two previous groups. This animal served as the normal control.

Samples of cecal tissue were removed from the 7 animals described above, and were fixed overnight at 4°C using 10% buffered formalin. The fixed tissues were paraffin-embedded, sectioned, and mounted on glass microscope slides. The tissue sections were then stained using hematoxylin and eosin (H and E stain), and were examined by light microscopy.

The tissues obtained from the two 24 hr. animals which received mixtures containing either 10µg or 30µg of toxin A and anti-recombinant toxin A IgY were indistinguishable from the normal control, both in terms of gross pathology, as well as at the microscopic level. These observations provide further evidence for the ability of anti-recombinant toxin A IgY to effectively neutralize the *in vivo* enterotoxin activity of C. difficile toxin A, and thus its ability to prevent acute or lasting toxin A-induced pathology.

In contrast, the tissues from the two 24 hr. animals which received the toxin A + pre-immune IgY mixtures demonstrated significant pathology. In both of these groups, the mucosal layer was observed to be less organized than in the normal control tissue. The cytoplasm of the epithelial cells had a vacuolated appearance, and gaps were present between the epithelium and the underlying cell layers. The lamina propria was largely absent. Intestinal villi and crypts were significantly diminished, and appeared to have been overgrown by a planar layer of epithelial cells and fibroblasts. Therefore, although these animals overtly appeared to recover from the acute symptoms of toxin A intoxication, lasting pathologic alterations to the cecal mucosa had occurred.

The tissues obtained from the two acute animals which received mixtures of toxin A and pre-immune IgY demonstrated the most significant pathology. At the gross pathological level, both animals were observed to have severely distended ceca which were filled with watery, diarrhea-like material. At the microscopic level, the animal that was given the mixture containing 10µg of toxin A and pre-immune IgY was found to have a mucosal layer which had a ragged, damaged appearance, and a disorganized, compacted quality. The crypts were largely absent, and numerous breaks in the epithelium had occurred. There was also an influx of erythrocytes into spaces between the epithelial layer and the underlying tissue. The animal which had received the mixture containing 30µg of toxin A and pre-immune IgY demonstrated the most severe pathology. The cecal tissue of this animal had an appearance very similar to that observed in animals which had died from C. difficile disease. Widespread destruction of the mucosa was noted, and the epithelial layer had sloughed. Hemorrhagic areas containing large numbers of erythrocytes were very prevalent. All semblance of normal tissue architecture was absent from this specimen. In terms of the presentation of pathologic events, this in vivo hamster model of toxin Aintoxication correlates very closely with the pathologic consequences of C. difficile disease in hamsters. The results presented in this Example demonstrate that while antirecombinant toxin A (Interval 6) IgY is capable of only partially neutralizing the cytotoxic activity of C. difficile toxin A, the same antibody effectively neutralizes 100% of the in vivo enterotoxin activity of the toxin. While it is not intended that this invention be limited to this mechanism, this may be due to the cytotoxicity and enterotoxicity of C. difficile Toxin A as two separate and distinct biological functions.

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EXAMPLE 15

In Vivo Neutralization Of C. Difficile Toxin A By Antibodies Against Recombinant Toxin A Polypeptides

The ability of avian antibodies directed against the recombinant *C. difficile* toxin A fragment 1870-2680 (as expressed by pMA1870-2680; see Example 13) to neutralize the enterotoxic activity of toxin A was demonstrated in Example 14. The ability of avian antibodies (IgYs) directed against other recombinant toxin A epitopes to neutralize native toxin A *in vivo* was next evaluated. This example involved: (a) the preparation of IgYs against recombinant toxin A polypeptides; (b) *in vivo* protection of hamsters against toxin A by treatment with anti-recombinant toxin A IgYs and (c) quantification of specific antibody concentration in CTA and Interval 6 IgY PEG preparations.

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The nucleotide sequence of the coding region of the entire toxin A protein is listed in SEQ ID NO:5. The amino acid sequence of the entire toxin A protein is listed in SEQ ID NO:6. The amino acid sequence consisting of amino acid residues 1870 through 2680 of toxin A is listed in SEQ ID NO:7. The amino acid sequence consisting of amino acid residues 1870 through 1960 of toxin A is listed in SEQ ID NO:8.

a) Preparation Of IgY's Against Recombinant Toxin A Polypeptides

Eggs were collected from Leghorn hens which have been immunized with recombinant *C. difficile* toxin A polypeptide fragments encompassing the entire toxin A protein. The polypeptide fragments used as immunogens were: 1) pMA 1870-2680 (Interval 6), 2) pPA 1100-1450 (Interval 4), and 3) a mixture of fragments consisting of pMA 30-300 (Interval 1), pMA 300-660 (Interval 2), pMA 660-1100 (Interval 3) and pMA 1450-1870 (Interval 5). This mixture of immunogens is referred to as Interval 1235. The location of each interval within the toxin A molecule is shown in

Figure 15A. In Figure 15A, the following abbreviations are used: pP refers to the pET23 vector (New England BioLabs); pM refers to the pMALTM-c vector (New England BioLabs); A refers to toxin A; the numbers refer to the amino acid interval expressed in the clone. (For example, the designation pMA30-300 indicates that the recombinant clone encodes amino acids 30-300 of toxin A and the vector used was pMALTM-c).

The recombinant proteins were generated as described in Example 11. The IgYs were extracted and solubilized in 0.1M carbonate buffer pH 9.5 for oral administration as described in Example 14(a). The IgY reactivities against each individual recombinant interval was evaluated by ELISA as described in Example 13(c).

b) In Vivo Protection Of Hamsters Against Toxin A By Treatment With Anti-Recombinant Toxin A Antibodies

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The ability of antibodies raised against recombinant toxin A polypeptides to provide in vivo protection against the enterotoxic activity of toxin A was examined in the hamster model system. This assay was performed as described in Example 14(b). Briefly, for each 40-50 gram female Golden Syrian hamster (Charles River), 1 ml of IgY 4X (i.e., resuspended in 1/4 of the original yolk volume) PEG prep against Interval 6, Interval 4 or Interval 1235 was mixed with 30 µg (LD₁₀₀ oral dose) of C. difficile toxin A (Tech Lab). Preimmune IgY mixed with toxin A served as a negative control. Antibodies raised against C. difficile toxoid A (Example 8) mixed with toxin A (CTA) served as a positive control. The mixture was incubated for 1 hour at 37°C then orally administered to lightly etherized hamsters using an 18G feeding needle. The animals were then observed for the onset of diarrhea and death for a period of approximately 24 hours. The results are shown in Table 18.

TABLE 18
Study Outcome After 24 Hours

Treatment group	Healthy	Diarrhea ²	Dead'
Preimmune	0	0	7
CTA	5	0	0
Interval 6	- 6	1	0 .
Interval 4	0	1	6
Interval 1235	0	0	7

- Animal shows no sign of illness.
- Animal developed diarrhea, but did not die.
- 3 Animal developed diarrhea and died.

Pre-treatment of toxin A with IgYs against Interval 6 prevented diarrhea in 6 of 7 hamsters and completely prevented death in all 7. In contrast, as with preimmune IgY, IgYs against Interval 4 and Interval 1235 had no effect on the onset of diarrhea and death in the hamsters.

c) Quantification Of Specific Antibody Concentration In CTA And Interval 6 IgY PEG Preparations

To determine the purity of IgY PEG preparations, an aliquot of a pMA1870-2680 (Interval 6) IgY PEG preparation was chromatographed using HPLC and a KW-803 sizing column (Shodex). The resulting profile of absorbance at 280 nm is shown in Figure 16. The single large peak corresponds to the predicted MW of IgY. Integration of the area under the single large peak showed that greater than 95% of the protein eluted from the column was present in this single peak. This result demonstrated that the majority (>95%) of the material absorbing at 280 nm in the PEG preparation corresponds to IgY. Therefore, absorbance at 280 nm can be used to determine the total antibody concentration in PEG preparations.

To determine the concentration of Interval 6-specific antibodies (expressed as percent of total antibody) within the CTA and pMA1870-2680 (Interval 6) PEG preparations, defined quantities of these antibody preparations were affinity purified on

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a pPA1870-2680(H) (shown schematically in Figure 15B) affinity column and the specific antibodies were quantified. In Figure 15B the following abbreviations are used: pP refers to the pET23 vector (New England BioLabs); pM refers to the pMALTM-c vector (New England BioLabs); pG refers to the pGEX vector (Pharmacia); pB refers to the PinPointTM Xa vector (Promega); A refers to toxin A; the numbers refer to the amino acid interval expressed in the clone. The solid black ovals represent the MBP; the hatched ovals represent glutathione S-transferase; the hatched circles represent the biotin tag; and HHH represents the poly-histidine tag.

An affinity column containing recombinant toxin A repeat protein was made as follows. Four ml of PBS-washed Actigel resin (Sterogene) was coupled with 5-10 mg of pPA1870-2680 inclusion body protein [prepared as described in Example (17) and dialyzed into PBS] in a 15 ml tube (Falcon) containing 1/10 final volume Ald-coupling solution (1 M sodium cyanoborohydride). Aliquots of the supernatant from the coupling reactions, before and after coupling, were assessed by Coomassie staining of 7.5% SDS-PAGE gels. Based upon protein band intensities, greater than 6 mg of recombinant protein was coupled to the resin. The resin was poured into a 10 ml column (BioRad), washed extensively with PBS, pre-eluted with 4 M guanidine-HCl (in 10 mM Tris-HCl, pH 8.0; 0.005% thimerosal) and re-equilibrated with PBS. The column was stored at 4°C.

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Aliquots of a pMA1870-2680 (Interval 6) or a CTA IgY polyclonal antibody preparation (PEG prep) were affinity purified on the above affinity column as follows. The column was attached to an UV monitor (ISCO) and washed with PBS. For pMA1870-2680 IgY purification, a 2X PEG prep (filter sterilized using a 0.45 µ filter; approximately 500 mg total IgY) was applied. The column was washed with PBS until the baseline was re-established (the column flow-through was saved), washed with BBSTween to elute nonspecifically binding antibodies and re-equilibrated with PBS. Bound antibody was eluted from the column in 4 M guanidine-HCl (in 10 mM Tris-HCl, pH 8.0; 0.005% thimerosal). The entire elution peak was collected in a 15 ml tube (Falcon). The column was re-equilibrated and the column eluate was re-chromatographed as described above. The antibody preparation was quantified by UV

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absorbance (the elution buffer was used to zero the spectrophotometer). Total purified antibody was approximately 9 mg and 1 mg from the first and second chromatography passes, respectively. The low yield from the second pass indicated that most specific antibodies were removed by the first round of chromatography. The estimated percentage of Interval 6 specific antibodies in the pMA1870-2680 PEG prep is approximately 2%.

The percentage of Interval 6 specific antibodies in the CTA PEG prep was determined (utilizing the same column and methodology described above) to be approximately 0.5% of total IgY.

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A 4X PEG prep contains approximately 20 mg/ml IgY. Thus in b) above, approximately 400 μg specific antibody in the Interval 6 PEG prep neutralized 30 μg toxin A *in vivo*.

EXAMPLE 16

In Vivo Treatment Of C. difficile Disease In Hamsters By Recombinant Interval 6 Antibodies

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The ability of antibodies directed against recombinant Interval 6 of toxin A to protect hamsters in vivo from C. difficile disease was examined. This example involved: (a) prophylactic treatment of C. difficile disease and (b) therapeutic treatment of C. difficile disease.

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a) Prophylactic Treatment Of C. difficile Disease

This experiment was performed as described in Example 9(b). Three groups each consisting of 7 female 100 gram Syrian hamsters (Charles River) were prophylactically treated with either preimmune IgYs, IgYs against native toxin A and B [CTAB; see Example 8 (a) and (b)] or IgYs against Interval 6. IgYs were prepared as 4X PEG preparations as described in Example 9(a).

The animals were orally dosed 3 times daily, roughly at 4 hour intervals, for 12 days with 1 ml antibody preparations diluted in Ensure. Using estimates of specific antibody concentration from Example 15(c), each dose of the Interval 6 antibody prep contained approximately 400 µg of specific antibody. On day 2 each hamster was predisposed to C. difficile infection by the oral administration of 3.0 mg of Clindamycin-HCl (Sigma) in 1 ml of water. On day 3 the hamsters were orally challenged with 1 ml of C. difficile inoculum strain ATCC 43596 in sterile saline containing approximately 100 organisms. The animals were then observed for the onset of diarrhea and subsequent death during the treatment period. The results are shown in Table 19.

TABLE 19
Lethality After 12 Days Of Treatment

Treatment Group	Number Animals Alive	Number Animals Dead
Preimmune	0	7
CTAB	6	1
Interval 6	7	. 0

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Treatment of hamsters with orally-administered IgYs against Interval 6 successfully protected 7 out of 7 (100%) of the animals from C. difficile disease. One of the hamsters in this group presented with diarrhea which subsequently resolved during the course of treatment. As shown previously in Example 9, antibodies to native toxin A and toxin B were highly protective. In this Example, 6 out of 7 animals survived in the CTAB treatment group. All of the hamsters treated with preimmune sera came down with diarrhea and died. The survivors in both the CTAB and Interval 6 groups remained healthy throughout a 12 day post-treatment period. In particular, 6 out of 7 Interval 6-treated hamsters survived at least 2 weeks after termination of treatment which suggests that these antibodies provide a long-lasting cure. These results represent the first demonstration that antibodies generated against a recombinant region of toxin A can prevent CDAD when administered passively to animals. These results also indicate that antibodies raised against Interval 6 alone may

be sufficient to protect animals from C. difficile disease when administered prophylactically.

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Previously others had raised antibodies against toxin A by actively immunizing hamsters against a recombinant polypeptide located within the Interval 6 region [Lyerly, D.M., et al. (1990) Curr. Microbiol. 21:29]. Figure 17 shows schematically the location of the Lyerly, et al. intra-Interval 6 recombinant protein (cloned into the pUC vector) in comparison with the complete Interval 6 construct (pMA1870-2680) used herein to generate neutralizing antibodies directed against toxin A. In Figure 17, the solid black oval represents the MBP which is fused to the toxin A Interval 6 in pMA1870-2680.

The Lyerly, et al. antibodies (intra-Interval 6) were only able to partially protect hamsters against C. difficile infection in terms of survival (4 out of 8 animals survived) and furthermore, these antibodies did not prevent diarrhea in any of the animals. Additionally, animals treated with the intra-Interval 6 antibodies [Lyerly, et al. (1990), supra] died when treatment was removed.

In contrast, the experiment shown above demonstrates that passive administration of anti-Interval 6 antibodies prevented diarrhea in 6 out of 7 animals and completely prevented death due to CDAD. Furthermore, as discussed above, passive administration of the anti-Interval 6 antibodies provides a long lasting cure (i.e., treatment could be withdrawn without incident).

b) Therapeutic Treatment Of C. difficile Disease: In Vivo
Treatment Of An Established C. difficile Infection In
Hamsters With Recombinant Interval 6 Antibodies

The ability of antibodies against recombinant interval 6 of toxin A to therapeutically treat C. difficile disease was examined. The experiment was performed essentially as described in Example 10(b). Three groups, each containing seven to eight female Golden Syrian hamsters (100 g each; Charles River) were treated with either preimmune IgY, IgYs against native toxin A and toxin B (CTAB) and IgYs

against Interval 6. The antibodies were prepared as described above as 4X PEG preparations.

The hamsters were first predisposed to *C. difficile* infection with a 3 mg dose of Clindamycin-HCl (Sigma) administered orally in 1 ml of water. Approximately 24 hrs later, the animals were orally challenged with 1 ml of *C. difficile* strain ATCC 43596 in sterile saline containing approximately 200 organisms. One day after infection, the presence of toxin A and B was determined in the feces of the hamsters using a commercial immunoassay kit (Cytoclone A+B EPA, Cambridge Biotech) to verify establishment of infection. Four members of each group were randomly selected and tested. Feces from an uninfected hamster was tested as a negative control. All infected animals tested positive for the presence of toxin according to the manufacturer's procedure. The initiation of treatment then started approximately 24 hr post-infection.

The animals were dosed daily at roughly 4 hr intervals with 1 ml antibody preparation diluted in Ensure® (Ross Labs). The amount of specific antibodies given per dose (determined by affinity purification) was estimated to be about 400 µg of anti-Interval 6 IgY (for animals in the Interval 6 group) and 100 µg and 70 µg of antitoxin A (Interval 6-specific) and anti-toxin B (Interval 3-specific; see Example 19), respectively, for the CTAB preparation. The animals were treated for 9 days and then observed for an additional 4 days for the presence of diarrhea and death. The results indicating the number of survivors and the number of dead 4 days post-infection are shown in Table 20.

TABLE 20
In vivo Therapeutic Treatment With Interval 6 Antibodies

Treatment Group	Number Animals Alive	Number Animals Dead
Preimmune	4	3
CTAB	8	0
Interval 6	8	0

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Antibodies directed against both Interval 6 and CTAB successfully prevented death from C. difficile when therapeutically administered 24 hr after infection. This result is significant since many investigators begin therapeutic treatment of hamsters with existing drugs (e.g., vancomycin, phenelfamycins, tiacumicins, etc.) 8 hr post-infection [Swanson, et al. (1991) Antimicrobial Agents and Chemotherapy 35:1108 and (1989) J. Antibiotics 42:94].

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Forty-two percent of hamsters treated with preimmune IgY died from CDAD. While the anti-Interval 6 antibodies prevented death in the treated hamsters, they did not eliminate all symptoms of CDAD as 3 animals presented with slight diarrhea. In addition, one CTAB-treated and one preimmune-treated animal also had diarrhea 14 days post-infection. These results indicate that anti-Interval 6 antibodies provide an effective means of therapy for CDAD.

EXAMPLE 17

Induction Of Toxin A Neutralizing Antibodies Requires Soluble Interval 6 Protein

As shown in Examples 11(d) and 15, expression of recombinant proteins in *E. coli* may result in the production of either soluble or insoluble protein. If insoluble protein is produced, the recombinant protein is solubilized prior to immunization of animals. To determine whether, one or both of the soluble or insoluble recombinant proteins could be used to generate neutralizing antibodies to toxin A, the following experiment was performed. This example involved a) expression of the toxin A repeats and subfragments of these repeats in *E. coli* using a variety of expression vectors; b) identification of recombinant toxin A repeats and sub-regions to which neutralizing antibodies bind; and c) determination of the neutralization ability of antibodies raised against soluble and insoluble toxin A repeat immunogen.

Expression Of The Toxin A Repeats And Subfragments Of These Repeats In E. coli Using A Variety Of Expression Vectors

The Interval 6 immunogen utilized in Examples 15 and 16 was the pMA1870-2680 protein, in which the toxin A repeats are expressed as a soluble fusion protein with the MBP (described in Example 11). Interestingly, expression of this region (from the Spel site to the end of the repeats, see Figure 15B) in three other expression constructs, as either native (pPA1870-2680), poly-His tagged [pPA1870-2680 (H)] or biotin-tagged (pBA1870-2680) proteins resulted in completely insoluble protein upon induction of the bacterial host (see Figure 15B). The host strain BL21 (Novagen) was used for expression of pBA1870-2680 and host strain BL21(DE3) (Novagen) was used for expression of pPA1870-2680 and pPA1870-2680(H). These insoluble proteins accumulated to high levels in inclusion bodies. Expression of recombinant plasmids in E. coli host cells grown in 2X YT medium was performed as described [Williams, et al. (1995), supra].

As summarized in Figure 15B, expression of fragments of the toxin A repeats (as either N-terminal *SpeI-EcoRI* fragments, or C-terminal *EcoRI*-end fragments) also yielded high levels of insoluble protein using pGEX (pGA1870-2190), PinPointTM-Xa (pBA1870-2190 and pBA2250-2680) and pET expression systems (pPA1870-2190). The pGEX and pET expression systems are described in Example 11. The PinPointTM-Xa expression system drives the expression of fusion proteins in *E. coli*. Fusion proteins from PinPointTM-Xa vectors contain a biotin tag at the amino-terminal end and can be affinity purified SoftLinkTM Soft Release avidin resin (Promega) under mild denaturing conditions (5 mM biotin).

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The solubility of expressed proteins from the pPG1870-2190 and pPA1870-2190 expression constructs was determined after induction of recombinant protein expression under conditions reported to enhance protein solubility [These conditions comprise growth of the host at reduced temperature (30°C) and the utilization of high (1 mM IPTG) or low (0.1 mM IPTG) concentrations of inducer [Williams et al.

(1995), supra]. All expressed recombinant toxin A protein was insoluble under these conditions. Thus, expression of these fragments of the toxin A repeats in pET and pGEX expression vectors results in the production of insoluble recombinant protein even when the host cells are grown at reduced temperature and using lower concentrations of the inducer. Although expression of these fragments in pMal vectors yielded affinity purifiable soluble fusion protein, the protein was either predominantly insoluble (pMA1870-2190) or unstable (pMA2250-2650). Attempts to solubilize expressed protein from the pMA1870-2190 expression construct using reduced temperature or lower inducer concentration (as described above) did not improve fusion protein solubility.

Collectively, these results demonstrate that expression of the toxin A repeat region in *E. coli* results in the production of insoluble recombinant protein, when expressed as either large (aa 1870-2680) or small (aa 1870-2190 or aa 2250-2680) fragments, in a variety of expression vectors (native or poly-his tagged pET, pGEX or PinPointTM- Xa vectors), utilizing growth conditions shown to enhance protein solubility. The exception to this rule were fusions with the MBP, which enhanced protein solubility, either partially (pMA1870-2190) or fully (pMA1870-2680).

b) Identification Of Recombinant Toxin A Repeats And Sub-Regions To Which Neutralizing Antibodies Bind

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Toxin A repeat regions to which neutralizing antibodies bind were identified by utilizing recombinant toxin A repeat region proteins expressed as soluble or insoluble proteins to deplete protective antibodies from a polyclonal pool of antibodies against native C. difficile toxin A. An in vivo assay was developed to evaluate proteins for the ability to bind neutralizing antibodies.

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The rational for this assay is as follows. Recombinant proteins were first premixed with antibodies against native toxin A (CTA antibody; generated in Example 8) and allowed to react. Subsequently, C. difficile toxin A was added at a concentration lethal to hamsters and the mixture was administered to hamsters via IP injection. If

the recombinant protein contains neutralizing epitopes, the CTA antibodies would lose their ability to bind toxin A resulting in diarrhea and/or death of the hamsters.

The assay was performed as follows. The lethal dose of toxin A when delivered orally to nine 40 to 50 g Golden Syrian hamsters (Sasco) was determined to be 10 to 30 µg. The PEG-purified CTA antibody preparation was diluted to 0.5X concentration (i.e., the antibodies were diluted at twice the original yolk volume) in 0.1 M carbonate buffer, pH 9.5. The antibodies were diluted in carbonate buffer to protect them from acid degradation in the stomach. The concentration of 0.5X was used because it was found to be the lowest effective concentration against toxin A. The concentration of Interval 6-specific antibodies in the 0.5X CTA prep was estimated to be 10-15 µg/ml (estimated using the method described in Example 15).

The inclusion body preparation [insoluble Interval 6 protein; pPA1870-2680(H)] and the soluble Interval 6 protein [pMA1870-2680; see Figure 15] were both compared for their ability to bind to neutralizing antibodies against *C. difficile* toxin A (CTA). Specifically, 1 to 2 mg of recombinant protein was mixed with 5 ml of a 0.5X CTA antibody prep (estimated to contain 60-70 µg of Interval 6-specific antibody). After incubation for 1 hr at 37°C, CTA (Tech Lab) at a final concentration of 30 µg/ml was added and incubated for another 1 hr at 37°C. One ml of this mixture containing 30 µg of toxin A (and 10-15 µg of Interval 6-specific antibody) was administered orally to 40-50 g Golden Syrian hamsters (Sasco). Recombinant proteins that result in the loss of neutralizing capacity of the CTA antibody would indicate that those proteins contain neutralizing epitopes. Preimmune and CTA antibodies (both at 0.5X) without the addition of any recombinant protein served as negative and positive controls, respectively.

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Two other inclusion body preparations, both expressed as insoluble products in the pET vector, were tested; one containing a different insert (toxin B fragment) other than Interval 6 called pPB1850-2070 (see Figure 18) which serves as a control for insoluble Interval 6, the other was a truncated version of the Interval 6 region called pPA1870-2190 (see Figure 15B). The results of this experiment are shown in Table 21.

TABLE 21

Binding Of Neutralizing Antibodies By Soluble Interval 6 Protein Study Outcome After 24 Hours

Treatment Group!	Healthy ²	Diarrhea ³	Dead*
Preimmune Ab	0	3	2
CTA Ab	4	1	0
CTA Ab + Int 6 (soluble)	1	2	2
CTA Ab + Int 6 (insoluble)	5	0	0
CTA Ab + pPB1850-2070	5	0	0
CTA Ab + pPA1870-2190	. 5	0.	0

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- C. difficile toxin A (CTA) was added to each group.
- Animals showed no signs of illness.
- Animals developed diarrhea but did not die.
- Animals developed diarrhea and died.

Preimmune antibody was ineffective against toxin A, while anti-CTA antibodies at a dilute 0.5X concentration almost completely protected the hamsters against the enterotoxic effects of CTA. The addition of recombinant proteins pPB1850-2070 or pPA1870-2190 to the anti-CTA antibody had no effect upon its protective ability, indicating that these recombinant proteins do not bind to neutralizing antibodies. On the other hand, recombinant Interval 6 protein was able to bind to neutralizing anti-CTA antibodies and neutralized the *in vivo* protective effect of the anti-CTA antibodies. Four out of five animals in the group treated with anti-CTA antibodies mixed with soluble Interval 6 protein exhibited toxin associated toxicity (diarrhea and death). Moreover, the results showed that Interval 6 protein must be expressed as a soluble product in order for it to bind to neutralizing anti-CTA antibodies since the addition of insoluble Interval 6 protein had no effect on the neutralizing capacity of the CTA antibody prep.

c) Determination Of Neutralization Ability Of Antibodies
Raised Against Soluble And Insoluble Toxin A Repeat
Immunogen

To determine if neutralizing antibodies are induced against solubilized inclusion bodies, insoluble toxin A repeat protein was solubilized and specific antibodies were raised in chickens. Insoluble pPA1870-2680 protein was solubilized using the method described in Williams et al. (1995), supra. Briefly, induced cultures (500 ml) were pelleted by centrifugation at 3,000 X g for 10 min at 4°C. The cell pellets were resuspended thoroughly in 10 ml of inclusion body sonication buffer (25 mM HEPES pH 7.7, 100 mM KCl, 12.5 mM MgCl₂, 20% glycerol, 0.1% (v/v) Nonidet P-40, 1 mM DTT). The suspension was transferred to a 30 ml non-glass centrifuge tube. Five hundred µl of 10 mg/ml lysozyme was added and the tubes were incubated on ice for 30 min. The suspension was then frozen at -70°C for at least 1 hr. The suspension was thawed rapidly in a water bath at room temperature and then placed on ice. The suspension was then sonicated using at least eight 15 sec bursts of the microprobe (Branson Sonicator Model No. 450) with intermittent cooling on ice.

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The sonicated suspension was transferred to a 35 ml Oakridge tube and centrifuged at 6,000 X g for 10 min at 4°C to pellet the inclusion bodies. The pellet was washed 2 times by pipetting or vortexing in fresh, ice-cold RIPA buffer [0.1% SDS, 1% Triton X-100, 1% sodium deoxycholate in TBS (25 mM Tris-Cl pH 7.5, 150 mM NaCl)]. The inclusion bodies were recentrifuged after each wash. The inclusion bodies were dried and transferred using a small metal spatula to a 15 ml tube (Falcon). One ml of 10% SDS was added and the pellet was solubilized by gently pipetting the solution up and down using a 1 ml micropipettor. The solubilization was facilitated by heating the sample to 95°C when necessary.

Once the inclusion bodies were in solution, the samples were diluted with 9 volumes of PBS. The protein solutions were dialyzed overnight against a 100-fold volume of PBS containing 0.05% SDS at room temperature. The dialysis buffer was then changed to PBS containing 0.01% SDS and the samples were dialyzed for several hours to overnight at room temperature. The samples were stored at 4°C until used. Prior to further use, the samples were warmed to room temperature to allow any precipitated SDS to go back into solution.

The inclusion body preparation was used to immunize hens. The protein was dialyzed into PBS and emulsified with approximately equal volumes of CFA for the initial immunization or IFA for subsequent booster immunizations. On day zero, for each of the recombinant recombinant preparations, two egg laying white Leghorn hens were each injected at multiple sites (IM and SC) with 1 ml of recombinant protein-adjuvant mixture containing approximately 0.5-1.5 mg of recombinant protein. Booster immunizations of 1.0 mg were given of days 14 and day 28. Eggs were collected on day 32 and the antibody isolated using PEG as described in Example 14(a). High titers of toxin A specific antibodies were present (as assayed by ELISA, using the method described in Example 13). Titers were determined for both antibodies against recombinant polypeptides pPA1870-2680 and pMA1870-2680 and were found to be comparable at > 1:62,500.

Antibodies against soluble Interval 6 (pMA1870-2680) and insoluble Interval 6 [(inclusion body), pPA1870-2680] were tested for neutralizing ability against toxin A using the *in vivo* assay described in Example 15(b). Preimmune antibodies and antibodies against toxin A (CTA) served as negative and positive controls, respectively. The results are shown in Table 22.

Neutralization Of Toxin A By Antibodies Against
Soluble Interval 6 Protein Study Outcome After 24 Hours

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Antibody Treatment Group	Healthy ¹	Diarrhea ²	Dead ³
Preimmune	l	0	4
СТА	5	0	0
Interval 6 (Soluble)4	. 5	0	0
Interval 6 (Insoluble)	0	2	3

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- Animals showed no sign of illness.
- Animal developed diarrhea but did not die.
- Animal developed diarrhea and died.
- 400 μg/ml.

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Antibodies raised against native toxin A were protective while preimmune antibodies had little effect. As found using the *in vitro* CHO assay [described in

Example 8(d)] where antibodies raised against the soluble Interval 6 could partially neutralize the effects of toxin A, here they were able to completely neutralize toxin A in vivo. In contrast, the antibodies raised against the insoluble Interval 6 was unable to neutralize the effects of toxin A in vivo as shown above (Table 22) and in vitro as shown in the CHO assay [described in Example 8(d)].

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These results demonstrate that soluble toxin A repeat immunogen is necessary to induce the production of neutralizing antibodies in chickens, and that the generation of such soluble immunogen is obtained only with a specific expression vector (pMal) containing the toxin A region spanning aa 1870-2680. That is to say, insoluble protein that is subsequently solubilized does not result in a toxin A antigen that will elicit a neutralizing antibody.

EXAMPLE 18

Cloning And Expression Of The C. difficile Toxin B Gene

The toxin B gene has been cloned and sequenced; the amino acid sequence deduced from the cloned nucleotide sequence predicts a MW of 269.7 kD for toxin B [Barroso et al., Nucl. Acids Res. 18:4004 (1990)]. The nucleotide sequence of the coding region of the entire toxin B gene is listed in SEQ ID NO:9. The amino acid sequence of the entire toxin B protein is listed in SEQ ID NO:10. The amino acid sequence consisting of amino acid residues 1850 through 2360 of toxin B is listed in SEQ ID NO:11. The amino acid sequence consisting of amino acid residues 1750 through 2360 of toxin B is listed in SEQ ID NO:12.

Given the expense and difficulty of isolating native toxin B protein, it would be advantageous to use simple and inexpensive procaryotic expression systems to produce and purify high levels of recombinant toxin B protein for immunization purposes. Ideally, the isolated recombinant protein would be soluble in order to preserve native antigenicity, since solubilized inclusion body proteins often do not fold into native conformations. Indeed as shown in Example 17, neutralizing antibodies against recombinant toxin A were only obtained when soluble recombinant toxin A

polypeptides were used as the immunogen. To allow ease of purification, the recombinant protein should be expressed to levels greater than 1 mg/liter of *E. coli* culture.

To determine whether high levels of recombinant toxin B protein could be produced in E. coli, fragments of the toxin B gene were cloned into various prokaryotic expression vectors, and assessed for the ability to express recombinant toxin B protein in E. coli. This Example involved (a) cloning of the toxin B gene and (b) expression of the toxin B gene in E. coli.

a) Cloning Of The Toxin B Gene

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The toxin B gene was cloned using PCR amplification from C. difficile genomic DNA. Initially, the gene was cloned in two overlapping fragments, using primer pairs P5/P6 and P7/P8. The location of these primers along the toxin B gene is shown schematically in Figure 18. The sequence of each of these primers is:

P5: 5' TAGAAAAAATGGCAAATGT 3' (SEQ ID NO:11); P6: 5' TTTCATCTTGTA GAGTCAAAG 3' (SEQ ID NO:12); P7: 5' GATGCCACAAGATGATTTAGTG 3' (SEQ ID NO:13); and P8: 5' CTAATTGAGCTGTATCAGGATC 3' (SEQ ID NO:14).

Figure 18 also shows the location of the following primers along the toxin B gene: P9 which consists of the sequence 5' CGGAATTCCTAGAAAAAATGGCAA ATG 3' (SEQ ID NO:15); P10 which consists of the sequence 5' GCTCTAGAATGA CCATAAGCTAGCCA 3' (SEQ ID NO:16); P11 which consists of the sequence 5' CGGAATTCGAGTTGGTAGAAAGGTGGA 3' (SEQ ID NO:17); P13 which consists of the sequence 5' CGGAATTCGGTTATTATCTTAAGGATG 3' (SEQ ID NO:18); and P14 which consists of the sequence 5' CGGAATTCTTGATAACTGGAT TTGTGAC 3' (SEQ ID NO:19). The amino acid sequence consisting of amino acid residues 1852 through 2362 of toxin B is listed in SEQ ID NO:20. The amino acid sequence consisting of amino acid sequence consisting of amino acid residues 1755 through 2362 of toxin B is listed in SEQ ID NO:21.

Clostridium difficile VPI strain 10463 was obtained from the American Type Culture Collection (ATCC 43255) and grown under anaerobic conditions in brain-heart infusion medium (Becton Dickinson). High molecular-weight C. difficile DNA was isolated essentially as described [Wren and Tabaqchali (1987) J. Clin. Microbiol., 25:2402], except 1) 100 µg/ml proteinase K in 0.5% SDS was used to disrupt the bacteria and 2) cetytrimethylammonium bromide (CTAB) precipitation [as described by Ausubel et al., Eds., Current Protocols in Molecular Biology, Vol. 2 (1989) Current Protocols] was used to remove carbohydrates from the cleared lysate. Briefly, after disruption of the bacteria with proteinase K and SDS, the solution is adjusted to approximately 0.7 M NaCl by the addition of a 1/7 volume of 5M NaCl. A 1/10 volume of CTAB/NaCl (10% CTAB in 0.7 M NaCl) solution was added and the solution was mixed thoroughly and incubated 10 min at 65°C. An equal volume of chloroform/isoamyl alcohol (24:1) was added and the phases were thoroughly mixed. The organic and aqueous phases were separated by centrifugation in a microfuge for 5 min. The aqueous supernatant was removed and extracted with phenol/chloroform/ isoamyl alcohol (25:24:1). The phases were separated by centrifugation in a microfuge for 5 min. The supernatant was transferred to a fresh tube and the DNA was precipitated with isopropanol. The DNA precipitate was pelleted by brief centrifugation in a microfuge. The DNA pellet was washed with 70% ethanol to remove residual CTAB. The DNA pellet was then dried and redissolved in TE buffer (10 mM Tris-HCl pH8.0, 1 mM EDTA). The integrity and yield of genomic DNA was assessed by comparison with a serial dilution of uncut lambda DNA after electrophoresis on an agarose gel.

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Toxin B fragments were cloned by PCR utilizing a proofreading thermostable DNA polymerase [native *Pfu* polymerase (Stratagene)]. The high fidelity of this polymerase reduces the mutation problems associated with amplification by error prone polymerases (e.g., *Taq* polymerase). PCR amplification was performed using the PCR primer pairs P5 (SEQ ID NO:11) with P6 (SEQ ID NO:12) and P7 (SEQ ID NO:13) with P8 (SEQ ID NO:14) in 50 µl reactions containing 10 mM Tris-HCl pH8.3, 50

mM KCl, 1.5 mM MgCl₂, 200 μM of each dNTP, 0.2 μM each primer, and 50 ng C. difficile genomic DNA. Reactions were overlaid with 100 μl mineral oil, heated to 94°C for 4 min, 0.5μl native Pfu polymerase (Stratagene) was added, and the reactions were cycled 30 times at 94°C for 1 min, 50°C for 1 min, 72°C (2 min for each kb of sequence to be amplified), followed by 10 min at 72°C. Duplicate reactions were pooled, chloroform extracted, and ethanol precipitated. After washing in 70% ethanol, the pellets were resuspended in 50 μl TE buffer (10 mM Tris-HCl pH8.0, 1 mM EDTA).

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The P5/P6 amplification product was cloned into pUC19 as outlined below. 10µl aliquots of DNA were gel purified using the Prep-a-Gene kit (BioRad), and ligated to Smal restricted pUC19 vector. Recombinant clones were isolated and confirmed by restriction digestion using standard recombinant molecular biology techniques (Sambrook et al., 1989). Inserts from two independent isolates were identified in which the toxin B insert was oriented such that the vector BamHI and SacI sites were 5' and 3' oriented, respectively (pUCB10-1530). The insert-containing BamHI/SacI fragment was cloned into similarly cut pET23a-c vector DNA, and protein expression was induced in small scale cultures (5 ml) of identified clones.

Total protein extracts were isolated, resolved on SDS-PAGE gels, and toxin B protein identified by Western analysis utilizing a goat anti-toxin B affinity purified antibody (Tech Lab). Procedures for protein induction, SDS-PAGE, and Western blot analysis were performed as described in Williams et al. (1995), supra. In brief, 5 ml cultures of bacteria grown in 2XYT containing 100 µg/ml ampicillin containing the appropriate recombinant clone were induced to express recombinant protein by addition of IPTG to 1mM. The E. coli hosts used were: BL21(DE3) or BL21(DE3)LysS (Novagen) for pET plasmids.

Cultures were induced by the addition of IPTG to a final concentration of 1.0 mM when the cell density reached 0.5 OD₆₀₀, and induced protein was allowed to accumulate for two hrs after induction. Protein samples were prepared by pelleting 1 ml aliquots of bacteria by centrifugation (1 min in microfuge), and resuspension of the

pelleted bacteria in 150 μl of 2X SDS-PAGE sample buffer (0.125 mM Tris-HCl pH 6.8, 2 mM EDTA, 6% SDS, 20% glycerol, 0.025% bromophenol blue; β-mercaptoethanol is added to 5% before use). The samples were heated to 95°C for 5 min, then cooled and 5 or 10 μls loaded on 7.5% SDS-PAGE gels. High molecular weight protein markers (BioRad) were also loaded, to allow estimation of the MW of identified fusion proteins. After electrophoresis, protein was detected either generally by staining the gels with Coomassie Blue, or specifically, by blotting to nitrocellulose for Western blot detection of specific immunoreactive protein. The MW of induced toxin B reactive protein allowed the integrity of the toxin B reading frame to be determined.

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The pET23b recombinant (pPB10-1530) expressed high MW recombinant toxin B reactive protein, consistent with predicted values. This confirmed that frame terminating errors had not occurred during PCR amplification. A pET23b expression clone containing the 10-1750aa interval of the toxin B gene was constructed, by fusion of the EcoRV-SpeI fragment of the P7/P8 amplification product to the P5-EcoRV interval of the P5/P6 amplification product (see Figure 18) in pPB10-1530. The integrity of this clone (pPB10-1750) was confirmed by restriction mapping, and Western blot detection of expressed recombinant toxin B protein. Levels of induced protein from both pPB10-1530 and pPB10-1750 were too low to facilitate purification of usable amounts of recombinant toxin B protein. The remaining 1750-2360 aa interval was directly cloned into pMAL or pET expression vectors from the P7/P8 amplification product as described below.

b) Expression Of The Toxin B Gene

i) Overview Of Expression Methodologies

proteins in E. coli. Native proteins were expressed in either the pET23a-c or pET16b expression vectors (Novagen). The pET23 vectors contain an extensive polylinker

sequence in all three reading frames (a-c vectors) followed by a C-terminal polyhistidine repeat. The pET16b vector contains a N-terminal poly-histidine sequence

Fragments of the toxin B gene were expressed as either native or fusion

immediately 5' to a small polylinker. The poly-histidine sequence binds to Ni-Chelate columns and allows affinity purification of tagged target proteins [Williams et al. (1995), supra]. These affinity tags are small (10 aa for pET16b, 6 aa for pET23) allowing the expression and affinity purification of native proteins with only limited amounts of foreign sequences.

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An N-terminal histidine-tagged derivative of pET16b containing an extensive cloning cassette was constructed to facilitate cloning of N-terminal poly-histidine tagged toxin B expressing constructs. This was accomplished by replacement of the promoter region of the pET23a and b vectors with that of the pET16b expression vector. Each vector was restricted with Bg/II and NdeI, and the reactions resolved on a 1.2 % agarose gel. The pET16b promoter region (contained in a 200 bp Bg/II-NdeI fragment) and the promoter-less pET23 a or b vectors were cut from the gel, mixed and Prep-A-Gene (BioRad) purified. The eluted DNA was ligated, and transformants screened for promoter replacement by NcoI digestion of purified plasmid DNA (the pET16b promoter contains this site, the pET23 promoter does not). These clones (denoted pETHisa or b) contain the pET16b promoter (consisting of a pT7-lac promoter, ribosome binding site and poly-histidine (10aa) sequence) fused at the NdeI site to the extensive pET23 polylinker.

All MBP fusion proteins were constructed and expressed in the pMAL[™]-c or pMAL[™]-p2 vectors (New England Biolabs) in which the protein of interest is expressed as a C-terminal fusion with MBP. All pET plasmids were expressed in either the BL21(DE3) or BL21(DE3)LysS expression hosts, while pMal plasmids were expressed in the BL21 host.

Large scale (500 mls to 1 liter) cultures of each recombinant were grown in 2X YT broth, induced, and soluble protein fractions were isolated as described [Williams, et al. (1995), supra]. The soluble protein extracts were affinity chromatographed to isolate recombinant fusion protein, as described [Williams et al., (1995) supra]. In brief, extracts containing tagged pET fusions were chromatographed on a nickel chelate column, and eluted using imidazole salts or low pH (pH 4.0) as described by

the distributor (Novagen or Qiagen). Extracts containing soluble pMAL fusion protein were prepared and chromatographed in PBS buffer over an amylose resin (New England Biolabs) column, and eluted with PBS containing 10 mM maltose as described [Williams et al. (1995), supra].

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ii) Overview Of Toxin B Expression

In both large expression constructs described in (a) above, only low level (i.e., less than 1 mg/liter of intact or nondegraded recombinant protein) expression of recombinant protein was detected. A number of expression constructs containing smaller fragments of the toxin B gene were then constructed, to determine if small regions of the gene can be expressed to high levels (i.e., greater than 1 mg/liter intact protein) without extensive protein degradation. All were constructed by in frame fusions of convenient toxin B restriction fragments to either the pMAL-c, pET23a-c, pET16b or pETHisa-b expression vectors, or by engineering restriction sites at specific locations using PCR amplification [using the same conditions described in (a) above]. In all cases, clones were verified by restriction mapping, and, where indicated, DNA sequencing.

Protein preparations from induced cultures of each of these constructs were analyzed, by SDS-PAGE, to estimate protein stability (Coomassie Blue staining) and immunoreactivity against anti-toxin B specific antiserum (Western analysis). Higher levels of intact (i.e., nondegraded), full length fusion proteins were observed with the smaller constructs as compared with the larger recombinants, and a series of expression constructs spanning the entire toxin B gene were constructed (Figures 18, 19 and 20 and Table 23).

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Constructs that expressed significant levels of recombinant toxin B protein (greater than 1 mg/liter intact recombinant protein) in E. coli were identified and a series of these clones that spans the toxin B gene are shown in Figure 19 and summarized in Table 23. These clones were utilized to isolate pure toxin B recombinant protein from the entire toxin B gene. Significant protein yields were

obtained from pMAL expression constructs spanning the entire toxin B gene, and yields of full length soluble fusion protein ranged from an estimated 1 mg/liter culture (pMB1100-1530) to greater than 20 mg/liter culture (pMB1750-2360).

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Representative purifications of MBP and poly-histidine-tagged toxin B recombinants are shown in Figures 21 and 22. Figure 21 shows a Coomassie Blue stained 7.5% SDS-PAGE gel on which various protein samples extracted from bacteria harboring pMB1850-2360 were electrophoresed. Samples were loaded as follows: Lane 1: protein extracted from uninduced culture; Lane 2: induced culture protein; Lane 3: total protein from induced culture after sonication; Lane 4: soluble protein; and Lane 5: eluted affinity purified protein. Figure 22 depicts the purification of recombinant proteins expressed in bacteria harboring either pPB1850-2360 (Lanes 1-3) or pPB1750-2360 (Lanes 4-6). Samples were loaded as follows: uninduced total protein (Lanes 1 and 4); induced total protein (Lanes 2 and 5); and eluted affinity purified protein (Lanes 3 and 6). The broad range molecular weight protein markers (BioRad) are shown in Lane 7.

Thus, although high level expression was not attained using large expression constructs from the toxin B gene, usable levels of recombinant protein were obtained by isolating induced protein from a series of smaller pMAL expression constructs that span the entire toxin B gene.

These results represent the first demonstration of the feasibility of expressing recombinant toxin B protein to high levels in *E. coli*. As well, expression of small regions of the putative ligand binding domain (repeat region) of toxin B as native protein yielded insoluble protein, while large constructs, or fusions to MBP were soluble (Figure 19), demonstrating that specific methodologies are necessary to produce soluble fusion protein from this interval.

iii) Clone Construction And Expression Details

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A portion of the toxin B gene containing the toxin B repeat region [amino acid residues 1852-2362 of toxin B (SEQ ID NO:20)] was isolated by PCR amplification of this interval of the toxin B gene from *C. difficile* genomic DNA. The sequence, and location within the toxin B gene, of the two PCR primers [P7 (SEQ ID NO:13) and P8 (SEQ ID NO:14)] used to amplify this region are shown in Figure 18.

DNA from the PCR amplification was purified by chloroform extraction and ethanol precipitation as described above. The DNA was restricted with *SpeI*, and the cleaved DNA was resolved by agarose gel electrophoresis. The restriction digestion with *SpeI* cleaved the 3.6 kb amplification product into a 1.8 kb doublet band. This doublet band was cut from the gel and mixed with appropriately cut, gel purified pMALc or pET23b vector. These vectors were prepared by digestion with *HindIII*, filling in the overhanging ends using the Klenow enzyme, and cleaving with *XbaI* (pMALc) or *NheI* (pET23b). The gel purified DNA fragments were purified using the Prep-A-Gene kit (BioRad) and the DNA was ligated, transformed and putative recombinant clones analyzed by restriction mapping.

pET and pMal clones containing the toxin B repeat insert (aa interval 1750-2360 of toxin B) were verified by restriction mapping, using enzymes that cleaved specific sites within the toxin B region. In both cases fusion of the toxin B Spel site with either the compatible Xbal site (pMal) or compatible Nhel site (pET) is predicted to create an in frame fusion. This was confirmed in the case of the pMB1750-2360 clone by DNA sequencing of the clone junction and 5' end of the toxin B insert using a MBP specific primer (New England Biolabs). In the case of the pET construct, the fusion of the blunt ended toxin B 3' end to the filled HindIII site should create an inframe fusion with the C-terminal poly-histidine sequence in this vector. The pPB1750-2360 clone selected had lost, as predicted, the HindIII site at this clone junction; this eliminated the possibility that an additional adenosine residue was added to the 3' end of the PCR product by a terminal transferase activity of the Pfu polymerase, since

fusion of this adenosine residue to the filled *Hind*III site would regenerate the restriction site (and was observed in several clones).

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One liter cultures of each expression construct were grown, and fusion protein purified by affinity chromatography on either an amylose resin column (pMAL constructs; resin supplied by New England Biolabs) or Ni-chelate column (pET constructs; resin supplied by Qiagen or Novagen) as described [Williams et al. (1995), supra]. The integrity and purity of the fusion proteins were determined by Coomassie staining of SDS-PAGE protein gels as well as Western blot analysis with either an affinity purified goat polyclonal antiserum (Tech Lab), or a chicken polyclonal PEG prep, raised against the toxin B protein (CTB) as described above in Example 8. In both cases, affinity purification resulted in yields in excess of 20 mg protein per liter culture, of which greater than 90% was estimated to be full-length recombinant protein. It should be noted that the poly-histidine affinity tagged protein was released from the Qiagen Ni-NTA resin at low imidazole concentration (60 mM), necessitating the use of a 40 mM imidazole rather than a 60 mM imidazole wash step during purification.

A periplasmically secreted version of pMB1750-2360 was constructed by replacement of the promoter and MBP coding region of this construct with that from a related vector (pMALTM-p2; New England Biolabs) in which a signal sequence is present at the N-terminus of the MBP, such that fusion protein is exported. This was accomplished by substituting a *BgIII-EcoRV* promoter fragment from pMAL-p2 into pMB1750-2360. The yields of secreted, affinity purified protein (recovered from osmotic shock extracts as described by Riggs in *Current Protocols in Molecular Biology*, Vol. 2, Ausubel, *et al.*, Eds. (1989), Current Protocols, pp. 16.6.1 - 16.6.14] from this vector (pMBp1750-2360) were 6.5 mg/liter culture, of which 50% was estimated to be full-length fusion protein.

The interval was also expressed in two non-overlapping fragments. pMB1750-1970 was constructed by introduction of a frameshift into pMB1750-2360, by restriction with *Hind*III, filling in the overhanging ends and religation of the plasmid.

Recombinant clones were selected by loss of the *Hind*III site, and further restriction map analysis. Recombinant protein expression from this vector was more than 20 mg/liter of greater than 90% pure protein.

The complementary region was expressed in pMB1970-2360. This construct was created by removal of the 1750-1970 interval of pMB1750-2360. This was accomplished by restriction of this plasmid with *EcoRI* (in the pMalc polylinker 5' to the insert) and III, filling in the overhanging ends, and religation of the plasmid. The resultant plasmid, pMB1970-2360, was made using both intracellularly and secreted versions of the pMB1750-2360 vector.

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No fusion protein was secreted in the pMBp1970-2360 version, perhaps due to a conformational constraint that prevents export of the fusion protein. However, the intracellularly expressed vector produced greater than 40mg/liter of greater than 90% full-length fusion protein.

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Constructs to precisely express the toxin B repeats in either pMalc (pMB1850-2360) or pET16b (pPB1850-2360) were constructed as follows. The DNA interval including the toxin B repeats was PCR amplified as described above utilizing PCR primers P14 (SEQ ID NO:19) and P8 (SEQ ID NO:14). Primer P14 adds a *Eco*RI site immediately flanking the start of the toxin B repeats.

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The amplified fragment was cloned into the pT7 Blue T-vector (Novagen) and recombinant clones in which single copies of the PCR fragment were inserted in either orientation were selected (pT71850-2360) and confirmed by restriction mapping. The insert was excised from two appropriately oriented independently isolated pT71850-2360 plasmids, with *EcoRI* (5' end of repeats) and *PstI* (in the flanking polylinker of the vector), and cloned into *EcoRI/PstI* cleaved pMalc vector. The resulting construct (pMB1850-2360) was confirmed by restriction analysis, and yielded 20 mg/l of soluble fusion protein [greater than 90% intact (*i.e.*, nondegraded)] after affinity chromatography. Restriction of this plasmid with *Hind*III and religation of the vector resulted in the removal of the 1970-2360 interval. The resultant construct (pMB1850-1970) expressed greater than 70 mg/liter of 90% full length fusion protein.

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The pPB1850-2360 construct was made by cloning a EcoRI (filled with Klenow)-BamHI fragment from a pT71850-2360 vector (opposite orientation to that used in the pMB1850-2360 construction) into NdeI (filled)/BamHI cleaved pET16b vector. Yields of affinity purified soluble fusion protein were 15 mg/liter, of greater than 90% full length fusion protein.

Several smaller expression constructs from the 1750-2070 interval were also constructed in His-tagged pET vectors, but expression of these plasmids in the BL21 (DE3) host resulted in the production of high levels of mostly insoluble protein (see Table 23 and Figure 19). These constructs were made as follows.

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pPB1850-1970 was constructed by cloning a Bg/II-HindIII fragment of pPB1850-2360 into Bg/II/HindIII cleaved pET23b vector. pPB1850-2070 was constructed by cloning a Bg/II-PvuII fragment of pPB1850-2360 into Bg/II/HincII cleaved pET23b vector. pPB1750-1970(c) was constructed by removal of the internal HindIII fragment of a pPB1750-2360 vector in which the vector HindIII site was regenerated during cloning (presumably by the addition of an A residue to the amplified PCR product by terminal transferase activity of Pfu polymerase). The pPB1750-1970(n) construct was made by insertion of the insert containing the Ndel-HindIII fragment of pPB1750-2360 into identically cleaved pETHisb vector. All constructs were confirmed by restriction digestion.

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An expression construct that directs expression of the 10-470 aa interval of toxin B was constructed in the pMalc vector as follows. A NheI (a site 5' to the insert in the pET23 vector)-AfIII (filled) fragment of the toxin B gene from pPB10-1530 was cloned into XbaI (compatible with NheI)/HindIII (filled) pMalc vector. The integrity of the construct (pMB10-470) was verified by restriction mapping and DNA sequencing of the 5' clone junction using a MBP specific DNA primer (New England Biolabs). However, all expressed protein was degraded to the MBP monomer MW.

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A second construct spanning this interval (aa 10-470) was constructed by cloning the PCR amplification product from a reaction containing the P9 (SEQ ID NO:15) and P10 (SEQ ID NO:16) primers (Figure 18) into the pETHisa vector. This

was accomplished by cloning the PCR product as an EcoRI-blunt fragment into EcoRI-HincII restricted vector DNA; recombinant clones were verified by restriction mapping. Although this construct (pPB10-520) allowed expression and purification (utilizing the N-terminal polyhistidine affinity tag) of intact fusion protein, yields were estimated at less than 500 µg per liter culture.

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Higher yield of recombinant protein from this interval (aa 10-520) were obtained by expression of the interval in two overlapping clones. The 10-330aa interval was cloned in both pETHisa and pMalc vectors, using the BamHI-AfIII (filled) DNA fragment from pPB10-520. This fragment was cloned into BamHI-HindIII (filled) restricted pMalc or BamHI-HindII restricted pETHisa vector. Recombinant clones were verified by restriction mapping. High level expression of either insoluble (pET) or soluble (pMal) fusion protein was obtained. Total yields of fusion protein from the pMB10-330 construct (Figure 18) were 20 mg/liter culture, of which 10% was estimated to be full-length fusion protein. Although yields of this interval were higher and >90% full-length recombinant protein produced when expressed from the pET construct, the pMal fusion was utilized since the expressed protein was soluble and thus more likely to have the native conformation.

The pMB260-520 clone was constructed by cloning *EcoRI-XbaI* cleaved amplified DNA from a PCR reaction containing the P11 (SEQ ID NO:17) and P10 (SEQ ID NO:16) DNA primers (Figure 18) into similarly restricted pMalc vector. Yields of affinity purified protein were 10 mg/liter, of which approximately 50% was estimated to be full-length recombinant protein.

The aa510-1110 interval was expressed as described below. This entire interval was expressed as a pMal fusion by cloning the *NheI-HindIII* fragment of pUCB10-1530 into *XbaI-HindIII* cleaved pMalc vector. The integrity of the construct (pMB510-1110) was verified by restriction mapping and DNA sequencing of the 5' clone junction using a MBP specific DNA primer. The yield of affinity purified protein was 25 mg/liter culture, of which 5% was estimated to be full-length fusion protein (1 mg/liter).

To attempt to obtain higher yields, this region was expressed in two fragments (aa510-820, and 820-1110) in the pMalc vector. The pMB510-820 clone was constructed by insertion of a SacI (in the pMalc polylinker 5' to the insert)-HpaI DNA fragment from pMB510-1110 into SacI/StuI restricted pMalc vector. The pMB820-1110 vector was constructed by insertion of the HpaI-HindIII fragment of pUCB10-1530 into BamHI (filled)/HindIII cleaved pMalc vector. The integrity of these constructs were verified by restriction mapping and DNA sequencing of the 5' clone junction using a MBP specific DNA primer.

Recombinant protein expressed from the pMB510-820 vector was highly unstable. However, high levels (20 mg/liter) of >90% full-length fusion protein were obtained from the pMB820-1105 construct. The combination of partially degraded pMB510-1110 protein (enriched for the 510-820 interval) with the pMB820-1110 protein provides usable amounts of recombinant antigen from this interval.

The aal 100-1750 interval was expressed as described below. The entire interval was expressed in the pMalc vector from a construct in which the AccI(filled)-SpeI fragment of pPB10-1750 was inserted into StuI/XbaI (XbaI is compatible with SpeI; StuI and filled AccI sites are both blunt ended) restricted pMalc. The integrity of this construct (pMB1100-1750) was verified by restriction mapping and DNA sequencing of the clone junction using a MBP specific DNA primer. Although 15 mg/liter of affinity purified protein was isolated from cells harboring this construct, the protein was greater than 99% degraded to MBP monomer size.

A smaller derivative of pMB1100-1750 was constructed by restriction of pMB1100-1750 with AfIII and SalI (in the pMalc polylinker 3' to the insert), filling in the overhanging ends, and religating the plasmid. The resultant clone (verified by restriction digestion and DNA sequencing) has deleted the aa1530-1750 interval, was designated pMB1100-1530. pMB1100-1530 expressed recombinant protein at a yield of greater than 40 mg/liter, of which 5% was estimated to be full-length fusion protein.

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Three constructs were made to express the remaining interval. Initially, a BspHI (filled)-SpeI fragment from pPB10-1750 was cloned into EcoRI(filled)/XbaI cleaved pMalc vector. The integrity of this construct (pMB1570-1750) was verified by restriction mapping and DNA sequencing of the 5' clone junction using a MBP specific DNA primer. Expression of recombinant protein from this plasmid was very low, approximately 3 mg affinity purified protein per liter, and most was degraded to MBP monomer size. This region was subsequently expressed from a PCR amplified DNA fragment. A PCR reaction utilizing primers P13 [SEQ ID NO:18; P13 was engineered to introduce an EcoRI site 5' to amplified toxin B sequences] and P8 (SEQ ID NO:14) was performed on C. difficile genomic DNA as described above. The amplified fragment was cleaved with EcoRI and SpeI, and cloned into EcoRI/XbaI cleaved pMalc vector. The resultant clone (pMB1530-1750) was verified by restriction map analysis, and recombinant protein was expressed and purified. The yield was greater than 20 mg protein per liter culture and it was estimated that 25% was fulllength fusion protein; this was a significantly higher yield than the original pMB1570-1750 clone. The insert of pMB1530-1750 (in a EcoRI-Sall fragment) was transferred to the pETHisa vector (EcoRI/XhoI cleaved, XhoI and SalI ends are compatible). No detectable fusion protein was purified on Ni-Chelate columns from soluble lysates of cells induced to express fusion protein from this construct.

20 TABLE 23
Summary Of Toxin B Expression Constructs*

Clone	Affinity Tag	Yield (mg/liter)	% Full Length
pPB10-1750	попе	low (estimated from Western blot hyb.)	?
pPB10-1530	none	low (as above)	?
pMB10-470	MBP	15mg/l	0%
pPB10-520	poly-his	0.5mg/l	20%
pPB10-330	poly-his	>20mg/l (insoluble)	90%
рМВ10-330	MBP	20mg/l	10%
pMB260-520	MBP	10mg/l	50%

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TABLE 23
Summary Of Toxin B Expression Constructs*

	Clone	Affinity Tag	Yield (mg/liter)	% Full Length
	pMB510-1110	MBP	25mg/l	5%
	pMB510-820	МВР	degraded (by Western blot hyb)	
	pMB820-1110	MBP	20mg/l	90%
	pMB1100-1750	МВР	15mg/l	0%
5	pMB1100-1530	МВР	40mg/l	5%
	pMB1570-1750	МВР	3mg/l	<5%
	pPB1530-1750	poly-his	no purified protein detected	?
	pMB1530-1750	MBP	20mg/l	25%
	pMB1750-2360	MBP	>20mg/l	>90%
10	pMBp1750-2360	МВР	6.5mg/l (secreted)	50%
	pPB1750-2360	poly-his .	>20mg/l	>90%
	pMB1750-1970	МВР	>20mg/l	>90%
	pMB1970-2360	МВР	40mg/l	>90%
	pMBp1970-2360	МВР	(no secretion)	NA
15	pMB1850-2360	МВР	20mg/l	>90%
	pPB1850-2360	poly-his	15mg/l	>90%
	pMB1850-1970	МВР	70mg/l	>90%
	pPB1850-1970	poly-his	>10mg/l (insoluble)	>90%
	pPB1850-2070	poly-his	>10mg/l (insoluble)	>90%
20	pPB1750-1970(c)	poly-his	>10mg/l (insoluble)	>90%
	pPB1750-1970(n)	poly-his	>10mg/l (insoluble)	>90%

Clones in italics are clones currently utilized to purify recombinant protein from each selected interval.

EXAMPLE 19

Identification, Purification And Induction Of Neutralizing
Antibodies Against Recombinant C. difficile Toxin B Protein

To determine whether recombinant toxin B polypeptide fragments can generate neutralizing antibodies, typically animals would first be immunized with recombinant proteins and anti-recombinant antibodies are generated. These anti-recombinant protein antibodies are then tested for neutralizing ability in vivo or in vitro. Depending on the immunogenic nature of the recombinant polypeptide, the generation of high-titer antibodies against that protein may take several months. To accelerate this process and identify which recombinant polypeptide(s) may be the best candidate to generate neutralizing antibodies, depletion studies were performed. Specifically, recombinant toxin B polypeptide were pre-screened by testing whether they have the ability to bind to protective antibodies from a CTB antibody preparation and hence deplete those antibodies of their neutralizing capacity. Those recombinant polypeptides found to bind CTB, were then utilized to generate neutralizing antibodies. This Example involved: a) identification of recombinant sub-regions within toxin B to which neutralizing antibodies bind; b) identification of toxin B sub-region specific antibodies that neutralize toxin B in vivo; and c) generation and evaluation of antibodies reactive to recombinant toxin B polypeptides.

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a) Identification Of Recombinant Sub-Regions Within Toxin B To Which Neutralizing Antibodies Bind

Sub-regions within toxin B to which neutralizing antibodies bind were identified by utilizing recombinant toxin B proteins to deplete protective antibodies from a polyclonal pool of antibodies against native C. difficile toxin B. An in vivo assay was developed to evaluate protein preparations for the ability to bind neutralizing

antibodies. Recombinant proteins were first pre-mixed with antibodies directed against native toxin B (CTB antibody; see Example 8) and allowed to react for one hour at 37°C. Subsequently, C. difficile toxin B (CTB; Tech Lab) was added at a concentration lethal to hamsters and incubated for another hour at 37°C. After incubation this mixture was injected intraperitoneally (IP) into hamsters. If the recombinant polypeptide contains neutralizing epitopes, the CTB antibodies will lose its ability to protect the hamsters against death from CTB. If partial or complete protection occurs with the CTB antibody-recombinant mixture, that recombinant contains only weak or non-neutralizing epitopes of toxin B. This assay was performed as follows.

Antibodies against CTB were generated in egg laying Leghorn hens as described in Example 8. The lethal dosage (LD 100) of *C. difficile* toxin B when delivered I.P. into 40g female Golden Syrian hamsters (Charles River) was determined to be 2.5 to 5 µg. Antibodies generated against CTB and purified by PEG precipitation could completely protect the hamsters at the I.P. dosage determined above. The minimal amount of CTB antibody needed to afford good protection against 5 µg of CTB when injected I.P. into hamsters was also determined (1X PEG prep). These experiments defined the parameters needed to test whether a given recombinant protein could deplete protective CTB antibodies.

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The cloned regions tested for neutralizing ability cover the entire toxin B gene and were designated as Intervals (INT) 1 through 5 (see Figure 19). Approximately equivalent final concentrations of each recombinant polypeptide were tested. The following recombinant polypeptides were used: 1) a mixture of intervals 1 and 2 (INT-1, 2); 2) a mixture of Intervals 4 and 5 (INT-4, 5) and 3) Interval 3 (INT-3). Recombinant proteins (each at about 1 mg total protein) were first preincubated with a final CTB antibody concentration of 1X [i.e., pellet dissolved in original yolk volume as described in Example 1(c)] in a final volume of 5 mls for 1 hour at 37°C. Twenty-

five μg of CTB (at a concentration of 5 μg/ml), enough CTB to kill 5 hamsters, was then added and the mixture was then incubated for 1 hour at 37°C. Five, 40g female hamsters (Charles River) in each treatment group were then each given 1 ml of the mixture I.P. using a tuberculin syringe with a 27 gauge needle. The results of this experiment are shown in Table 24.

TABLE 24

Binding Of Neutralizing Antibodies By INT 3 Protein

Treatment Group	Number Of Animals Alive	Number Of Animals Dead
CTB antibodies	3	2
CTB antibodies + INT1,2	3	2
CTB antibodies + INT4,5	3	2
CTB antibodies + INT 3	0	5

C. difficile toxin B (CTB) was added to each group.

As shown in Table 24, the addition of recombinant proteins from INT-1, 2 or INT-4, 5 had no effect on the *in vivo* protective ability of the CTB antibody preparation compared to the CTB antibody preparation alone. In contrast, INT-3 recombinant polypeptide was able to remove all of the toxin B neutralizing ability of the CTB antibodies as demonstrated by the death of all the hamsters in that group.

The above experiment was repeated, using two smaller expressed fragments (pMB 1750-1970 and pMB 1970-2360, see Figure 19) comprising the INT-3 domain to determine if that domain could be further subdivided into smaller neutralizing epitopes. In addition, full-length INT-3 polypeptide expressed as a nickel tagged protein (pPB1750-2360) was tested for neutralizing ability and compared to the original INT-3 expressed MBP fusion (pMB1750-2360). The results are shown in Table 25.

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TABLE 25

Removal Of Neutralizing Antibodies By Repeat Containing Proteins

Treatment Group!	Number Of Animals Alive	Number Of Animals Dead
CTB antibodies	5	0
CTB antibodies + pPB1750-2360	0	5
CTB antibodies + pMB1750-2360	0	5
CTB antibodies + pMB1970-2360	3	2
CTB antibodies + pMB1750-1970	2	3

C. difficile toxin B (CTB) was added to each group.

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The results summarized in Table 25 indicate that the smaller polypeptide fragments within the INT-3 domain, pMB1750-1970 and pMB1970-2360, partially lose the ability to bind to and remove neutralizing antibodies from the CTB antibody pool. These results demonstrate that the full length INT-3 polypeptide is required to completely deplete the CTB antibody pool of neutralizing antibodies. This experiment also shows that the neutralization epitope of INT-3 can be expressed in alternative vector systems and the results are independent of the vector utilized or the accompanying fusion partner.

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Other Interval 3 specific proteins were subsequently tested for the ability to remove neutralizing antibodies within the CTB antibody pool as described above. The Interval 3 specific proteins used in these studies are summarized in Figure 23. In Figure 23 the following abbreviations are used: pP refers to the pET23 vector; pM refers to the pMALc vector; B refers to toxin B; the numbers refer to the amino acid interval expressed in the clone. The solid black ovals represent the MBP; and HHH represents the poly-histidine tag.

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Only recombinant proteins comprising the entire toxin B repeat domain (pMB1750-2360, pPB1750-2360 and pPB1850-2360) can bind and completely remove neutralizing antibodies from the CTB antibody pool. Recombinant proteins comprising only a portion of the toxin B repeat domain were not capable of completely removing

neutralizing antibodies from the CTB antibody pool (pMB1750-1970 and pMB1970-2360 could partially remove neutralizing antibodies while pMB1850-1970 and pPB1850-2070 failed to remove any neutralizing antibodies from the CTB antibody pool).

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The above results demonstrate that only the complete ligand binding domain (repeat region) of the toxin B gene can bind and completely remove neutralizing antibodies from the CTB antibody pool. These results demonstrate that antibodies directed against the entire toxin B repeat region are necessary for *in vivo* toxin neutralization (see Figure 23; only the recombinant proteins expressed by the pMB1750-2360, pPB1750-2360 and pPB1850-2360 vectors are capable of completely removing the neutralizing antibodies from the CTB antibody pool).

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These results represent the first indication that the entire repeat region of toxin B would be necessary for the generation of antibodies capable of neutralizing toxin B, and that sub-regions may not be sufficient to generate maximal titers of neutralizing antibodies.

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b) Identification Of Toxin B Sub-Region Specific Antibodies That Neutralize Toxin B In Vivo

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To determine if antibodies directed against the toxin B repeat region are sufficient for neutralization, region specific antibodies within the CTB antibody preparation were affinity purified, and tested for *in vivo* neutralization. Affinity columns containing recombinant toxin B repeat proteins were made as described below. A separate affinity column was prepared using each of the following recombinant toxin B repeat proteins: pPB1750-2360, pPB1850-2360, pMB1750-1970 and pMB1970-2360.

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For each affinity column to be made, four ml of PBS-washed Actigel resin (Sterogene) was coupled overnight at room temperature with 5-10 mg of affinity purified recombinant protein (first extensively dialyzed into PBS) in 15 ml tubes (Falcon) containing a 1/10 final volume Ald-coupling solution (1 M sodium cyanoborohydride). Aliquots of the supernatants from the coupling reactions, before

and after coupling, were assessed by Coomassie staining of 7.5% SDS-PAGE gels. Based on protein band intensities, in all cases greater than 30% coupling efficiencies were estimated. The resins were poured into 10 ml columns (BioRad), washed extensively with PBS, pre-eluted with 4M guanidine-HCl (in 10 mM Tris-HCl, pH 8.0) and reequilibrated in PBS. The columns were stored at 4°C.

Aliquots of a CTB IgY polyclonal antibody preparation (PEG prep) were affinity purified on each of the four columns as described below. The columns were hooked to a UV monitor (ISCO), washed with PBS and 40 ml aliquots of a 2X PEG prep (filter sterilized using a 0.45 μ filter) were applied. The columns were washed with PBS until the baseline value was re-established. The columns were then washed with BBStween to elute nonspecifically binding antibodies, and reequilibrated with PBS. Bound antibody was eluted from the column in 4M guanidine-HCl (in 10mM Tris-HCl, pH8.0). The eluted antibody was immediately dialyzed against a 100-fold excess of PBS at 4°C for 2 hrs. The samples were then dialyzed extensively against at least 2 changes of PBS, and affinity purified antibody was collected and stored at 4°C. The antibody preparations were quantified by UV absorbance. The elution volumes were in the range of 4-8 ml. All affinity purified stocks contained similar total antibody concentrations, ranging from 0.25-0.35% of the total protein applied to the columns.

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The ability of the affinity purified antibody preparations to neutralize toxin B in vivo was determined using the assay outlined in a) above. Affinity purified antibody was diluted 1:1 in PBS before testing. The results are shown in Table 26.

In all cases similar levels of toxin neutralization was observed, such that lethality was delayed in all groups relative to preimmune controls. This result demonstrates that antibodies reactive to the repeat region of the toxin B gene are sufficient to neutralize toxin B *in vivo*. The hamsters will eventually die in all groups, but this death is maximally delayed with the CTB PEG prep antibodies. Thus neutralization with the affinity purified (AP) antibodies is not as complete as that observed with the CTB prep before affinity chromatography. This result may be due to loss of activity during guanidine denaturation (during the elution of the antibodies

from the affinity column) or the presence of antibodies specific to other regions of the toxin B gene that can contribute to toxin neutralization (present in the CTB PEG prep).

TABLE 26

Neutralization Of Toxin B By Affinity Purified Antibodies

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Treatment group*	Number Animals Alive	Number Animals Dead
Preimmune ¹	0	5
CTB ¹ ; 400 μg	5	0
CTB (AP on pPB1750-2360); ² 875 μg	5	0
CTB (AP on pMB1750-1970); ² 875 μg	5	0
CTB (AP on pMB1970-2360); ² 500 μg	5	0

C. difficile toxin B (CTB) (Tech Lab; at 5 μg/ml, 25 μg total) at lethal concentration to hamsters is added to antibody and incubated for one hour at 37°C. After incubation this mixture is injected intraperitoneally (IP) into hamsters. Each treatment group received toxin premixed with antibody raised against the indicated protein, as either: ¹4X antibody PEG prep or ²affinity purified (AP) antibody (from CTB PEG prep, on indicated columns). The amount of specific antibody in each prep is indicated; the amount is directly determined for affinity purified preps and is estimated for the 4X CTB as described in Example 15.

The numbers in each group represent numbers of hamsters dead or alive, 2 hr post IP administration of toxin/antibody mixture.

The observation that antibodies affinity purified against the non-overlapping pMB1750-1970 and pMB1970-2360 proteins neutralized toxin B raised the possibility that either 1) antibodies specific to repeat sub-regions are sufficient to neutralize toxin B or 2) sub-region specific proteins can bind most or all repeat specific antibodies present in the CTB polyclonal pool. This would likely be due to conformational similarities between repeats, since homology in the primary amino acid sequences between different repeats is in the range of only 25-75% [Eichel-Streiber, et al. (1992) Molec. Gen. Genetics 233:260]. These possibilities were tested by affinity chromatography.

The CTB PEG prep was sequentially depleted 2X on the pMB1750-1970 column; only a small elution peak was observed after the second chromatography,

indicating that most reactive antibodies were removed. This interval depleted CTB preparation was then chromatographed on the pPB1850-2360 column; no antibody bound to the column. The reactivity of the CTB and CTB (pMB1750-1970 depleted) preps to pPB1750-2360, pPB1850-2360, pMB1750-1970 and pMB1970-2360 proteins was then determined by ELISA using the protocol described in Example 13(c). Briefly, 96-well microtiter plates (Falcon, Pro-Bind Assay Plates) were coated with recombinant protein by adding 100 μl volumes of protein at 1-2 μg/ml in PBS containing 0.005% thimerosal to each well and incubating overnight at 4°C. The next morning, the coating suspensions were decanted and the wells were washed three times using PBS. In order to block non-specific binding sites, 100 µl of 1.0% BSA (Sigma) in PBS (blocking solution) was then added to each well, and the plates were incubated for 1 hr. at 37°C. The blocking solution was decanted and duplicate samples of 150 ul of diluted antibody was added to the first well of a dilution series. The initial testing serum dilution was (1/200 for CTB prep, (the concentration of depleted CTB was standardized by OD₂₈₀) in blocking solution containing 0.5% Tween 20, followed by 5-fold serial dilutions into this solution. This was accomplished by serially transferring 30 µl aliquots to 120 µl buffer, mixing, and repeating the dilution into a fresh well. After the final dilution, 30 µl was removed from the well such that all wells contained 120 µl final volume. A total of 5 such dilutions were performed (4 wells total). The plates were incubated for 1 hr at 37°C. Following this incubation, the serially diluted samples were decanted and the wells were washed three times using PBS containing 0.5% Tween 20 (PBST), followed by two 5 min washes using BBS-Tween and a final three washes using PBST. To each well, 100 µl of 1/1000 diluted secondary antibody [rabbit anti-chicken IgG alkaline phosphatase (Sigma) diluted in blocking solution containing 0.5% Tween 20] was added, and the plate was incubated 1 hr at 37°C. The conjugate solutions were decanted and the plates were washed 6 times in PBST, then once in 50 mM Na₂CO₃, 10 mM MgCl₂, pH 9.5. The plates were developed by the addition of 100 µl of a solution containing 1 mg/ml para-nitro phenyl phosphate (Sigma) dissolved in 50 mM Na₂CO₃, 10 mM MgCl₂, pH9.5 to each well. The plates were then incubated at room temperature in the dark for 5-45 min.

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The absorbency of each well was measured at 410 nm using a Dynatech MR 700 plate reader.

As predicted by the affinity chromatography results, depletion of the CTB prep on the pMB1750-1970 column removed all detectable reactivity to the pMB1970-2360 protein. The reciprocal purification of a CTB prep that was depleted on the pMB1970-2360 column yielded no bound antibody when chromatographed on the pMB1750-1970 column. These results demonstrate that all repeat reactive antibodies in the CTB polyclonal pool recognize a conserved structure that is present in non-overlapping repeats. Although it is possible that this conserved structure represents rare conserved linear epitopes, it appears more likely that the neutralizing antibodies recognize a specific protein conformation. This conclusion was also suggested by the results of Western blot hybridization analysis of CTB reactivity to these recombinant proteins.

Western blots of 7.5% SDS-PAGE gels, loaded and electrophoresed with defined quantities of each recombinant protein, were probed with the CTB polyclonal antibody preparation. The blots were prepared and developed with alkaline phosphatase as described in Example 3. The results are shown in Figure 24.

Figure 24 depicts a comparison of immunoreactivity of IgY antibody raised against either native or recombinant toxin B antigen. Equal amounts of pMB1750-1970 (lane 1), pMB1970-2360 (lane 2), pPB1850-2360 (lane 3) as well as a serial dilution of pPB1750-2360 (lanes 4-6 comprising 1X, 1/10X and 1/100X amounts, respectively) proteins were loaded in duplicate and resolved on a 7.5% SDS-PAGE gel. The gel was blotted and each half was hybridized with PEG prep IgY antibodies from chickens immunized with either native CTB or pPB1750-2360 protein. Note that the full-length pMB1750-1970 protein was identified only by antibodies reactive to the recombinant protein (arrows).

Although the CTB prep reacts with the pPB1750-2360, pPB1850-2360, and pMB1970-2360 proteins, no reactivity to the pMB1750-1970 protein was observed (Figure 24). Given that all repeat reactive antibodies can be bound by this protein during affinity chromatography, this result indicates that the protein cannot fold

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properly on Western blots. Since this eliminates all antibody reactivity, it is unlikely that the repeat reactive antibodies in the CTB prep recognize linear epitopes. This may indicate that in order to induce protective antibodies, recombinant toxin B protein will need to be properly folded.

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- c) Generation And Evaluation Of Antibodies Reactive To Recombinant Toxin B Polypeptides
 - i) Generation Of Antibodies Reactive To Recombinant Toxin B Proteins

Antibodies against recombinant proteins were generated in egg laying Leghorn hens as described in Example 13. Antibodies were raised [using Freunds adjuvant (Gibco) unless otherwise indicated] against the following recombinant proteins: 1) a mixture of Interval 1+2 proteins (see Figure 18); 2) a mixture of interval 4 and 5 proteins (see Figure 18); 3) pMB1970-2360 protein; 4) pPB1750-2360 protein; 5) pMB1750-2360; 6) pMB1750-2360 [Titermax adjuvant (Vaxcell)]; 7) pMB1750-2360 [Gerbu adjuvant (Biotech)]; 8) pMBp1750-2360 protein; 9) pPB1850-2360; and 10) pMB1850-2360.

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Chickens were boosted at least 3 times with recombinant protein until ELISA reactivity [using the protocol described in b) above with the exception that the plates were coated with pPB1750-2360 protein] of polyclonal PEG preps was at least equal to that of the CTB polyclonal antibody PEG prep. ELISA titers were determined for the PEG preps from all of the above immunogens and were found to be comparable ranging from 1:12500 to 1:62500. High titers were achieved in all cases except in 6) pMB1750-2360 in which strong titers were not observed using the Titermax adjuvant, and this preparation was not tested further.

ii) Evaluation Of Antibodies Reactive To Recombinant Proteins By Western Blot Hybridization

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Western blots of 7.5% SDS-PAGE gels, loaded and electrophoresed with defined quantities of recombinant protein (pMB1750-1970, pPB1850-2360, and pMB1970-2360 proteins and a serial dilution of the pPB1750-2360 to allow quantification of reactivity), were probed with the CTB, pPB1750-2360, pMB1750-2360 and pMB1970-2360 polyclonal antibody preparations (from chickens immunized using Freunds adjuvant). The blots were prepared and developed with alkaline phosphatase as described above in b).

As shown in Figure 24, the CTB and pMB1970-2360 preps reacted strongly with the pPB1750-2360, pPB1850-2360, and pMB1970-2360 proteins while the pPB1750-2360 and pMB1970-2360 (Gerbu) preparations reacted strongly with all four proteins. The Western blot reactivity of the pPB1750-2360 and pMB1970-2360 (Gerbu) preparations were equivalent to that of the CTB preparation, while reactivity of the pMB1970-2360 preparation was <10% that of the CTB prep. Despite equivalent ELISA reactivities only weak reactivity (approximately 1%) to the recombinant proteins were observed in PEG preps from two independent groups immunized with the pMB1750-2360 protein and one group immunized with the pMB1750-2360 preparation using Freunds adjuvant.

Affinity purification was utilized to determine if this difference in immunoreactivity by Western blot analysis reflects differing antibody titers. Fifty ml 2X PEG preparations from chickens immunized with either pMB1750-2360 or pMB1970-2360 protein were chromatographed on the pPB1750-2360 affinity column from b) above, as described. The yield of affinity purified antibody (% total protein in preparation) was equivalent to the yield obtained from a CTB PEG preparation in b) above. Thus, differences in Western reactivity reflect a qualitative difference in the antibody pools, rather than quantitative differences., These results demonstrate that certain recombinant proteins are more effective at generating high affinity antibodies (as assayed by Western blot hybridization).

iii) In Vivo Neutralization Of Toxin B Using Antibodies Reactive To Recombinant Protein

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The in vivo hamster model [described in Examples 9 and 14(b)] was utilized to assess the neutralizing ability of antibodies raised against recombinant toxin B proteins.

The results from three experiments are shown below in Tables 27-29.

The ability of each immunogen to neutralize toxin B in vivo has been compiled and is shown in Table 30. As predicted from the recombinant protein-CTB premix studies (Table 24) only antibodies to Interval 3 (1750-2366) and not the other regions of toxin B (i.e., intervals 1-5) are protective. Unexpectedly, antibodies generated to INT-3 region expressed in pMAL vector (pMB1750-2360 and pMpB1750-2360) using Freunds adjuvant were non-neutralizing. This observation is reproducible, since no neutralization was observed in two independent immunizations with pMB1750-2360 and one immunization with pMpB1750-2360. The fact that 5X quantities of affinity purified toxin B repeat specific antibodies from pMB1750-2360 PEG preps cannot neutralize toxin B while 1X quantities of affinity purified anti-CTB antibodies can (Table 28) demonstrates that the differential ability of CTB antibodies to neutralize toxin B is due to qualitative rather than quantitative differences in these antibody preparations. Only when this region was expressed in an alternative vector (pPB1750-2360) or using an alternative adjuvant with the pMB1750-2360 protein were neutralizing antibodies generated. Importantly, antibodies raised using Freunds adjuvant to pPB1850-2360, which contains a fragment that is only 100 amino acids smaller than recombinant pPB1750-2360, are unable to neutralize toxin B in vivo (Table 27); note also that the same vector is used for both pPB1850-2360 and pPB1750-2360.

TABLE 27

In Vivo Neutralization Of Toxin B

Treatment Group*	Number Animals Alive	Number Animals Dead ^b
Preimmune	0	5
СТВ	5	0
INT 1+2	0	5
INT 4+5	0	5
pMB1750-2360	0	5
рМВ1970-2360	0	5
pPB1750-2360	5	. 0

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C. difficile toxin B (CTB) (at 5 µg/ml; 25 µg total; Tech Lab) at lethal concentration to hamsters is added to antibody and incubated for one hour at 37°C. After incubation this mixture is injected intraperitoneally (IP) into hamsters. Each treatment group received toxin premixed with antibody raised against the indicated protein, as a 4X antibody PEG prep.

The numbers in each group represent numbers of hamsters dead or alive, 2 hours post IP administration of toxin/antibody mixture.

TABLE 28

In Vivo Neutralization Of Toxin B Using Affinity Purified Antibodies

Treatment Group	Number Animals Alive	Number Animals Dead
Preimmune(1)	0	5 '
CTB(I)	5	0
pPB1750-2360(1)	5	. 0
1.5 mg anti-pMB1750-2360(2)	l	. 4
1.5 mg anti-pMB1970-2360(2)	0	5
300 μg anti-CTB(2)	S	0

- C. difficile toxin B (CTB) (at 5 μg/ml; 25 μg total; Tech Lab) at lethal concentration to hamsters is added to antibody and incubated for one hour at 37°C. After incubation, 1 ml of this mixture is injected intraperitoneally (IP) into hamsters. Each treatment group received toxin premixed with antibody raised against the indicated protein, as either (1) 4X antibody PEG prep or (2) affinity purified antibody (on a pPB1750-2360 resin), either 1.5 mg/group (anti-pMB1750-2360 and anti-pMB1970-2360; used undiluted affinity purified antibody) or 350 μg/group (anti-CTB, repeat specific; used 1/5 diluted anti-CTB antibody).
 - The numbers in each group represent numbers of hamsters dead or alive, 2 hr post-IP administration of toxin/antibody mixture.

TABLE 29

Generation Of Neutralizing Antibodies Utilizing The Gerbu Adjuvant

Treatment Group*	Number Animals Alive	·
Preimmune	0	5 .
СТВ	5	0
pMB1970-2360	0	5
pMB1850-2360	0	5
pPB1850-2360	0	5
pMB1750-2360 (Gerbu adj)	5	0

- C. difficile toxin B (CTB) (Tech Lab) at lethal concentration to hamsters is added to antibody and incubated for one hour at 37°C. After incubation this mixture is injected intraperitoneally (IP) into hamsters. Each treatment group received toxin premixed with antibody raised against the indicated protein, as a 4X antibody PEG prep.
- The numbers in each group represent numbers of hamsters dead or alive, 2hrs post IP administration of toxin/antibody mixture.

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TABLE 30

In Vivo Neutralization Of Toxin B

Immunogen	Adjuvant	Tested Preparation!	Antigen Utilized For AP	In vivo
Preimmune	NA ¹	PEG	NA	no
CTB (native)	Titermax	PEG	NA	yes
CTB (native)	Titermax	AP	pPB1750-2360	yes .
CTB (native)	Titermax	AP	pPB1850-2360	yes
CTB (native)	Titermax	AP	pPB1750-1970	yes
CTB (native)	Titermax	AP	pPB1970-2360	yes
pMB1750-2360	Freunds	PEG	NA	no
pMB1750-2360	Freunds	AP	pPB1750-2360	no
pMB1750-2360	Gerbu	PEG	NA	yes
pMB1970-2360	Freunds	PEG	NA	по
pMB1970-2360	Freunds	AP	pPB1750-2360	no
pPB1750-2360	Freunds	PEG	NA	yes
pPB1850-2360	Freunds	PEG	NA	no
pMB1850-2360	Freunds	PEG	NA	по
INT 1+2	Freunds	PEG	NA	по
INT 4+5	Freunds	PEG	NA	по

- 20 Either PEG preparation (PEG) or affinity purified antibodies (AP).
 - Yes' denotes complete neutralization (0/5 dead) while 'no' denotes no neutralization (5/5 dead) of toxin B, 2 hours post-administration of mixture.
 - 'NA' denotes not applicable.

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The pPB1750-2360 antibody pool confers significant *in vivo* protection, equivalent to that obtained with the affinity purified CTB antibodies. This correlates with the observed high affinity of this antibody pool (relative to the pMB1750-2360 or pMB1970-2360 pools) as assayed by Western blot analysis (Figure 24). These results provide the first demonstration that *in vivo* neutralizing antibodies can be induced using recombinant toxin B protein as immunogen.

The failure of high concentrations of antibodies raised against the pMB1750-2360 protein (using Freunds adjuvant) to neutralize, while the use of Gerbu adjuvant and pMB1750-2360 protein generates a neutralizing response, demonstrates that conformation or presentation of this protein is essential for the induction of neutralizing antibodies. These results are consistent with the observation that the neutralizing antibodies produced when native CTB is used as an immunogen appear to recognize conformational epitopes [see section b) above]. This is the first demonstration that the conformation or presentation of recombinant toxin B protein is essential to generate high titers of neutralizing antibodies.

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EXAMPLE 20

Determination Of Quantitative And Qualitative

Differences Between pMB1750-2360, pMB1750-2360 (Gerbu)

Or pPB1750-2360 IgY Polyclonal Antibody Preparations

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In Example 19, it was demonstrated that toxin B neutralizing antibodies could be generated using specific recombinant toxin B proteins (pPB1750-2360) or specific adjuvants. Antibodies raised against pMB1750-2360 were capable of neutralizing the enterotoxin effect of toxin B when the recombinant protein was used to immunize hens in conjunction with the Gerbu adjuvant, but not when Freunds adjuvant was used. To determine the basis for these antigen and adjuvant restrictions, toxin B-specific antibodies present in the neutralizing and non-neutralizing PEG preparations were isolated by affinity chromatography and tested for qualitative or quantitative differences. The example involved a) purification of anti-toxin B specific antibodies from pMB1750-2360 and pPB1750-2360 PEG preparations and b) in vivo neutralization of toxin B using the affinity purified antibody.

a) Purification Of specific Antibodies From pMB1750-2360 And pPB1750-2360 PEG Preparations

To purify and determine the concentration of specific antibodies (expressed as the percent of total antibody) within the pPB1750-2360 (Freunds and Gerbu) and pPB1750-2360 PEG preparations, defined quantities of these antibody preparations were chromatographed on an affinity column containing the entire toxin B repeat region (pPB1750-2360). The amount of affinity purified antibody was then quantified.

An affinity column containing the recombinant toxin B repeat protein, pPB1750-2360, was made as follows. Four ml of PBS-washed Actigel resin (Sterogene) was coupled with 5 mg of pPB1750-2360 affinity purified protein (dialyzed into PBS; estimated to be greater than 95% full length fusion protein) in a 15 ml tube (Falcon) containing 1/10 final volume Ald-coupling solution (1M sodium cyanoborohydride). Aliquots of the supernatant from the coupling reactions, before and after coupling, were assessed by Coomassie staining of 7.5% SDS-PAGE gels. Based on protein band intensities, greater than 95% (approximately 5 mg) of recombinant protein was coupled to the resin. The coupled resin was poured into a 10 ml column (BioRad), washed extensively with PBS, pre-eluted with 4M guanidine-HCl (in 10 mM Tris-HCl, pH 8.0; 0.005% thimerosal) and re-equilibrated in PBS and stored at 4°C.

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Aliquots of pMB1750-2360, pMB1750-2360 (Gerbu) or pPB1750-2360 IgY polyclonal antibody preparations (PEG preps) were affinity purified on the above column as follows. The column was attached to an UV monitor (ISCO), and washed with PBS. Forty ml aliquots of 2X PEG preps (filter sterilized using a 0.45 μ filter and quantified by OD₂₈₀ before chromatography) was applied. The column was washed with PBS until the baseline was re-established (the column flow-through was saved), washed with BBSTween to elute nonspecifically binding antibodies and reequilibrated with PBS. Bound antibody was eluted from the column in 4M guanidine-HCl (in 10 mM Tris-HCL, pH 8.0, 0.005% thimerosal) and the entire elution peak collected in a 15 ml tube (Falcon). The column was re-equilibrated, and the column eluate re-chromatographed as described above. The antibody preparations were

quantified by UV absorbance (the elution buffer was used to zero the spectrophotometer). Approximately 10 fold higher concentrations of total purified antibody was obtained upon elution of the first chromatography pass relative to the second pass. The low yield from the second chromatography pass indicated that most of the specific antibodies were removed by the first round of chromatography.

Pools of affinity purified specific antibodies were prepared by dialysis of the column elutes after the first column chromatography pass for the pMB1750-2360, pMB1750-2360 (Gerbu) or pPB1750-2360 IgY polyclonal antibody preparations. The elutes were collected on ice and immediately dialyzed against a 100-fold volume of PBS at 4°C for 2 hrs. The samples were then dialyzed against 3 changes of a 65-fold volume of PBS at 4°C. Dialysis was performed for a minimum of 8 hrs per change of PBS. The dialyzed samples were collected, centrifuged to remove insoluble debris, quantified by OD₂₈₀, and stored at 4°C.

The percentage of toxin B repeat-specific antibodies present in each preparation was determined using the quantifications of antibody yields from the first column pass (amount of specific antibody recovered after first pass/total protein loaded). The yield of repeat-specific affinity purified antibody (expressed as the percent of total protein in the preparation) in: 1) the pMB1750-2360 PEG prep was approximately 0.5%, 2) the pMB1750-2360 (Gerbu) prep was approximately 2.3%, and 3) the pPB1750-2360 prep was approximately 0.4%. Purification of a CTB IgY polyclonal antibody preparation on the same column demonstrated that the concentration of toxin B repeat specific antibodies in the CTB preparation was 0.35%.

These results demonstrate that 1) the use of Gerbu adjuvant enhanced the titer of specific antibody produced against the pMB1750-2360 protein 5-fold relative to immunization using Freunds adjuvant, and 2) the differences seen in the *in vivo* neutralization ability of the pMB1750-2360 (not neutralizing) and pPB1750-2360 (neutralizing) and CTB (neutralizing) PEG preps seen in Example 19 was not due to differences in the titers of repeat-specific antibodies in the three preparations because the titer of repeat-specific antibody was similar for all three preps; therefore the differing ability of the three antibody preparations to neutralize toxin B must reflect

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qualitative differences in the induced toxin B repeat-specific antibodies. To confirm that qualitative differences exist between antibodies raised in hens immunized with different recombinant proteins and/or different adjuvants, the same amount of affinity purified anti-toxin B repeat (aa 1870-2360 of toxin B) antibodies from the different preparations was administered to hamsters using the *in vivo* hamster model as described below.

b) In vivo Neutralization Of Toxin B Using Affinity Purified Antibody

The *in vivo* hamster model was utilized to assess the neutralizing ability of the affinity purified antibodies raised against recombinant toxin B proteins purified in (a) above. As well, a 4X IgY PEG preparation from a second independent immunization utilizing the pPB1750-2360 antigen with Freunds adjuvant was tested for *in vivo* neutralization. The results are shown in Table 31.

The results shown in Table 31 demonstrate that:

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- as shown in Example 19 and reproduced here, 1.5 mg of affinity purified antibody from pMB1750-2360 immunized hens using Freunds adjuvant does not neutralize toxin B *in vivo*. However, 300 μg of affinity purified antibody from similarly immunized hens utilizing Gerbu adjuvant demonstrated complete neutralization of toxin B *in vivo*. This demonstrates that Gerbu adjuvant, in addition to enhancing the titer of antibodies reactive to the pMB1750-2360 antigen relative to Freunds adjuvant (demonstrated in (a) above), also enhances the yield of neutralizing antibodies to this antigen, greater than 5 fold.
- 2) Complete in vivo neutralization of toxin B was observed with 1.5 mg of affinity purified antibody from hens immunized with pPB1750-2360 antigen, but not with pMB1750-2360 antigen, when Freunds adjuvant was used. This demonstrates, using standardized toxin B repeat-specific antibody concentrations, that neutralizing antibodies were induced when

pPB1750-2360 but not pMB1750-2360 was used as the antigen with Freunds adjuvant.

Complete in vivo neutralization was observed with 300 μg of pMB1750-2360 (Gerbu) antibody, but not with 300 μg of pPB1750-2360 (Freunds) antibody. Thus the pMB1750-2360 (Gerbu) antibody has a higher titer of neutralizing antibodies than the pPB1750-2360 (Freunds) antibody.

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- 4) Complete neutralization of toxin B was observed using 300 μg of CTB antibody [affinity purified (AP)] but not 100 μg CTB antibody (AP or PEG prep). This demonstrates that greater than 100 μg of toxin B repeat-specific antibody (anti-CTB) is necessary to neutralize 25 μg toxin B in vivo in this assay, and that affinity purified antibodies specific to the toxin B repeat interval neutralize toxin B as effectively as the PEP prep of IgY raised against the entire CTB protein (shown in this assay).
 - As was observed with the initial pPB1750-2360 (IgY) PEG preparation (Example 19), complete neutralization was observed with a IgY PEG preparation isolated from a second independent group of pPB1750-2360 (Freunds) immunized hens. This demonstrates that neutralizing antibodies are reproducibly produced when hens are immunized with pPB1750-2360 protein utilizing Freunds adjuvant.

TABLE 31 In vivo Neutralization Of Toxin B Using Affinity Purified Antibodies

Treatment Group*	Number Animals Alive	Number Animals Deadb
Preimmune ¹	0	5
CTB (300 μg) ²	5	0
CTB (100 μg) ²	1	4
pMB1750-2360 (G) (5 mg) ²	5	0
pMB1750-2360 (G) (1.5 mg) ²	. 5	0
pMB1750-2360 (G) (300 μg) ²	· 5	0
pMB1750-2360 (F) (1.5 mg) ²	0	5
pPB1750-2360 (F) (1.5 mg) ²	5	0
pPB1750-2360 (F) (300 μg) ²	1	4
CTB (100 μg) ³	2	3
pPB1750-2360 (F) (500 μg) ¹	5	0

15 C. difficile toxin B (CTB) (Tech Lab) at lethal concentration to hamsters (25 µg) was added to the antibody (amount of specific antibody is indicated) and incubated for one hour at 37°C. After incubation, this mixture was injected IP into hamsters (1/5 total mix injected per hamster). Each treatment group received toxin premixed with antibody raised against the indicated protein (G=gerbu adjuvant, F=Freunds adjuvant). 20 indicates the antibody was a 4X IgY PEG prep; indicates the antibody was affinity purified on a pPB1850-2360 resin; and 3 indicates that the antibody was a 1X IgY PEG prep.

> The numbers in each group represent numbers of hamsters dead or alive, 2 hrs post IP administration of toxin/antibody mixture.

EXAMPLE 21

Diagnostic Enzyme Immunoassays For C. difficile Toxins A And B

The ability of the recombinant toxin proteins and antibodies raised against these recombinant proteins (described in the above examples) to form the basis of diagnostic assays for the detection of clostridial toxin in a sample was examined. Two immunoassay formats were tested to quantitatively detect C. difficile toxin A and toxin B from a biological specimen. The first format involved a competitive assay in which

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a fixed amount of recombinant toxin A or B was immobilized on a solid support (e.g., microtiter plate wells) followed by the addition of a toxin-containing biological specimen mixed with affinity-purified or PEG fractionated antibodies against recombinant toxin A or B. If toxin is present in a specimen, this toxin will compete with the immobilized recombinant toxin protein for binding to the anti-recombinant antibody thereby reducing the signal obtained following the addition of a reporter reagent. The reporter reagent detects the presence of antibody bound to the immobilized toxin protein.

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In the second format, a sandwich immunoassay was developed using affinity-purified antibodies to recombinant toxin A and B. The affinity-purified antibodies to recombinant toxin A and B were used to coat microtiter wells instead of the recombinant polypeptides (as was done in the competitive assay format). Biological samples containing toxin A or B were then added to the wells followed by the addition of a reporter reagent to detect the presence of bound toxin in the well.

a) Competitive Immunoassay For The Detection Of C. difficile Toxin

Recombinant toxin A or B was attached to a solid support by coating 96 well microtiter plates with the toxin protein at a concentration of 1µg/ml in PBS. The plates were incubated overnight at 2-8°C. The following morning, the coating solutions were removed and the remaining protein binding sites on the wells were blocked by filling each well with a PBS solution containing 0.5% BSA and 0.05% Tween-20. Native C. difficile toxin A or B (Tech Lab) was diluted to 4 µg/ml in stool extracts from healthy Syrian hamsters (Sasco). The stool extracts were made by placing fecal pellets in a 15 ml centrifuge tube; PBS was added at 2 ml/pellet and the tube was vortexed to create a uniform suspension. The tube was then centrifuged at 2000 rpm for 5 min at room temperature. The supernatant was removed; this comprises the stool extract. Fifty µl of the hamster stool extract was pipetted into each well of the microtiter plates to serve as the diluent for serial dilutions of the 4 µg/ml toxin samples. One hundred µl of the toxin samples at 4 µg/ml was pipetted

into the first row of wells in the microtiter plate, and 50 µl aliquots were removed and diluted serially down the plate in duplicate. An equal volume of affinity purified anti-recombinant toxin antibodies [1 ng/well of anti-pMA1870-2680 antibody was used for the detection of toxin A; 0.5 ng/well of anti-pMB1750-2360(Gerbu) was used for the detection of toxin B] were added to appropriate wells, and the plates were incubated at room temperature for 2 hours with gentle agitation. Wells serving as negative control contained antibody but no native toxin to compete for binding.

Unbound toxin and antibody were removed by washing the plates 3 to 5 times with PBS containing 0.05% Tween-20. Following the wash step, 100 µl of rabbit anti-chicken IgG antibody conjugated to alkaline phosphatase (Sigma) was added to each well and the plates were incubated for 2 hours at room temperature. The plates were then washed as before to remove unbound secondary antibody. Freshly prepared alkaline phosphatase substrate (1 mg/ml p-nitrophenyl phosphate (Sigma) in 50 mM Na₂CO₃, pH 9.5; 10 mM MgCl₂) was added to each well. Once sufficient color developed, the plates were read on a Dynatech MR700 microtiter plate reader using a 410 nm filter.

The results are summarized in Tables 32 and 33. For the results shown in Table 32, the wells were coated with recombinant toxin A protein (pMA1870-2680). The amount of native toxin A added (present as an addition to solubilized hamster stool) to a given well is indicated (0 to 200 ng). Antibody raised against the recombinant toxin A protein, pMA1870-2680, was affinity purified on the an affinity column containing pPA1870-2680 (described in Example 20). As shown in Table 32, the recombinant toxin A protein and affinity-purified antitoxin can be used for the basis of a competitive immunoassay for the detection of toxin A in biological samples.

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Similar results were obtained using the recombinant toxin B, pPB1750-2360, and antibodies raised against pMB1750-2360(Gerbu). For the results shown in Table 33, the wells were coated with recombinant toxin B protein (pPB1750-2360). The amount of native toxin B added (present as an addition to solubilized hamster stool) to a given well is indicated (0 to 200 ng). Antibody raised against the recombinant toxin B protein, pMB1750-2360(Gerbu), was affinity purified on the an

affinity column containing pPB1850-2360 (described in Example 20). As shown in Table 33, the recombinant toxin B protein and affinity-purified antitoxin can be used for the basis of a competitive immunoassay for the detection of toxin B in biological samples.

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In this competition assay, the reduction is considered significant over the background levels at all points; therefore the assay can be used to detect samples containing less than 12.5 ng toxin A/well and as little as 50-100 ng toxin B/well.

TABLE 32

Competitive Inhibition Of Anti-C. difficile Toxin A By Native Toxin A

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ng Toxin A/Well	OD ₄₁₀ Readout
200	0.176
100	0.253
50	0.240
25	0.259
12.5	0.309
6.25	0.367
3.125	0.417
0	0.590

TABLE 33

Competitive Inhibition Of Anti-C. difficile Toxin B By Native Toxin B

ng Toxin B/Well	OD ₄₁₀ Readout
200	0.392
100	0.566
50	0.607
25	0.778
12.5	0.970
6.25	0.902
3.125	1.040
0	1.055

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These competitive inhibition assays demonstrate that native C. difficile toxins and recombinant C. difficile toxin proteins can compete for binding to antibodies raised against recombinant C. difficile toxins demonstrating that these anti-recombinant toxin antibodies provide effective diagnostic reagents.

Sandwich Immunoassay For The Detection Of C. difficile Toxin

Affinity-purified antibodies against recombinant toxin A or toxin B were immobilized to 96 well microtiter plates as follows. The wells were passively coated overnight at 4°C with affinity purified antibodies raised against either pMA1870-2680 (toxin A) or pMB1750-2360(Gerbu) (toxin B). The antibodies were affinity purified as described in Example 20. The antibodies were used at a concentration of 1 µg/ml and 100 µl was added to each microtiter well. The wells were then blocked with 200 µl of 0.5% BSA in PBS for 2 hours at room temperature and the blocking solution was then decanted. Stool samples from healthy Syrian hamsters were resuspended in PBS, pH 7.4 (2 ml PBS/stool pellet was used to resuspend the pellets and the sample was centrifuged as described above). The stool suspension was then spiked with native C. difficile toxin A or B (Tech Lab) at 4 µg/ml. The stool suspensions

containing toxin (either toxin A or toxin B) were then serially diluted two-fold in stool suspension without toxin and 50 µl was added in duplicate to the coated microtiter wells. Wells containing stool suspension without toxin served as the negative control.

The plates were incubated for 2 hours at room temperature and then were washed three times with PBS. One hundred µl of either goat anti-native toxin A or goat anti-native toxin B (Tech Lab) diluted 1:1000 in PBS containing 1% BSA and 0.05% Tween 20 was added to each well. The plates were incubated for another 2 hours at room temperature. The plates were then washed as before and 100 µl of alkaline phosphatase-conjugated rabbit anti-goat IgG (Cappel, Durham, N.C.) was added at a dilution of 1:1000. The plates were incubated for another 2 hours at room temperature. The plates were washed as before then developed by the addition of 100 µl/well of a substrate solution containing 1 mg/ml p-nitrophenyl phosphate (Sigma) in 50 mM Na₂CO₃, pH 9.5; 10 mM MgCl₂. The absorbance of each well was measured using a plate reader (Dynatech) at 410 nm. The assay results are shown in Tables 34 and 35.

TABLE 34

C. difficile Toxin A Detection In Stool Using Affinity-Purified Antibodies Against Toxin A

	OD ₄₁₀ Readout
200	0.9
100	0.8
50	0.73
25	0.71
12.5	0.59
6.25	0.421
0	0

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TABLE 35

C. difficile Toxin B Detection In Stool Using Affinity-Purified Antibodies Against Toxin B

ng Toxin B/Well	OD ₄₁₀ Readout
200	1.2
. 100	0.973
50	0.887
25	0.846
12.5	0.651
6.25	0.431
0	0.004

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The results shown in Tables 34 and 35 show that antibodies raised against recombinant toxin A and toxin B fragments can be used to detect the presence of C. difficile toxin in stool samples. These antibodies form the basis for a sensitive sandwich immunoassay which is capable of detecting as little as 6.25 ng of either toxin A or B in a 50 µl stool sample. As shown above in Tables 34 and 35, the background for this sandwich immunoassay is extremely low; therefore, the sensitivity of this assay is much lower than 6.25 ng toxin/well. It is likely that toxin levels of 0.5 to 1.0 pg/well could be detected by this assay.

The results shown above in Tables 32-35 demonstrate clear utility of the recombinant reagents in *C. difficile* toxin detection systems.

EXAMPLE 22

Construction And Expression Of C. botulinum C Fragment Fusion Proteins

The C. botulinum type A neurotoxin gene has been cloned and sequenced [Thompson, et al., Eur. J. Biochem. 189:73 (1990)]. The nucleotide sequence of the toxin gene is available from the EMBL/GenBank sequence data banks under the accession number X52066; the nucleotide sequence of the coding region is listed in SEQ ID NO:27. The amino acid sequence of the C. botulinum type A neurotoxin is

listed in SEQ ID NO:28. The type A neurotoxin gene is synthesized as a single polypeptide chain which is processed to form a dimer composed of a light and a heavy chain linked via disulfide bonds. The 50 kD carboxy-terminal portion of the heavy chain is referred to as the C fragment or the H_C domain.

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Previous attempts by others to express polypeptides comprising the C fragment of C. botulinum type A toxin as a native polypeptide (e.g., not as a fusion protein) in E. coli have been unsuccessful [H.F. LaPenotiere, et al. in Botulinum and Tetanus Neurotoxins, DasGupta, Ed., Plenum Press, New York (1993), pp. 463-466]. Expression of the C fragment as a fusion with the E. coli MBP was reported to result in the production of insoluble protein (H.F. LaPenotiere, et al., supra).

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In order to produce soluble recombinant C fragment proteins in *E. coli*, fusion proteins comprising a synthetic C fragment gene derived from the *C. botulinum* type A toxin and either a portion of the *C. difficile* toxin protein or the MBP were constructed. This example involved a) the construction of plasmids encoding C fragment fusion proteins and b) expression of *C. botulinum* C fragment fusion proteins in *E. coli*.

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a) Construction Of Plasmids Encoding C Fragment Fusion Proteins

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In Example 11, it was demonstrated that the *C. difficile* toxin A repeat domain can be efficiently expressed and purified in *E. coli* as either native (expressed in the pET 23a vector in clone pPA1870-2680) or fusion (expressed in the pMALc vector as a fusion with the *E. coli* MBP in clone pMA1870-2680) proteins. Fusion proteins comprising a fusion between the MBP, portions of the *C. difficile* toxin A repeat domain (shown to be expressed as a soluble fusion protein) and the C fragment of the *C. botulinum* type A toxin were constructed. A fusion protein comprising the C fragment of the *C. botulinum* type A toxin and the MBP was also constructed.

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Figure 25 provides a schematic representation of the botulinal fusion proteins along with the donor constructs containing the *C. difficile* toxin A sequences or *C.*

botulinum C fragment sequences which were used to generate the botulinal fusion proteins. In Figure 25, the solid boxes represent C. difficile toxin A gene sequences, the open boxes represent C. botulinum C fragment sequences and the solid black ovals represent the E. coli MBP. When the name for a restriction enzyme appears inside parenthesis, this indicates that the restriction site was destroyed during construction. An asterisk appearing with the name for a restriction enzyme indicates that this restriction site was recreated at the cloning junction.

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In Figure 25, a restriction map of the pMA1870-2680 and pPA1100-2680 constructs (described in Example 11) which contain sequences derived from the C. difficile toxin A repeat domain are shown; these constructs were used as the source of C. difficile toxin A gene sequences for the construction of plasmids encoding fusions between the C. botulinum C fragment gene and the C. difficile toxin A gene. The pMA1870-2680 expression construct expresses high levels of soluble, intact fusion protein (20 mg/liter culture) which can be affinity purified on an amylose column (purification described in Example 11d).

The pAlterBot construct (Figure 25) was used as the source of *C. botulinum* C fragment gene sequences for the botulinal fusion proteins. pAlterBot was obtained from J. Middlebrook and R. Lemley at the U.S. Department of Defense. pAlterBot contains a synthetic *C. botulinum* C fragment inserted in to the pALTER-1® vector (Promega). This synthetic C fragment gene encodes the same amino acids as does the naturally occurring C fragment gene. The naturally occurring C fragment sequences, like most clostridial genes, are extremely A/T rich (Thompson *et al.*, *supra*). This high A/T content creates expression difficulties in *E. coli* and yeast due to altered codon usage frequency and fortuitous polyadenylation sites, respectively. In order to improve the expression of C fragment proteins in *E. coli*, a synthetic version of the gene was created in which the non-preferred codons were replaced with preferred codons.

The nucleotide sequence of the *C. botulinum* C fragment gene sequences contained within pAlterBot is listed in SEQ ID NO:22. The first six nucleotides

(ATGGCT) encode a methionine and alanine residue, respectively. These two amino acids result from the insertion of the *C. botulinum* C fragment sequences into the pALTER® vector and provide the initiator methionine residue. The amino acid sequence of the *C. botulinum* C fragment encoded by the sequences contained within pAlterBot is listed in SEQ ID NO:23. The first two amino acids (Met Ala) are encoded by vector-derived sequences. From the third amino acid residue onward (Arg), the amino acid sequence is identical to that found in the *C. botulinum* type A toxin gene.

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The pMA1870-2680, pPA1100-2680 and pAlterBot constructs were used as progenitor plasmids to make expression constructs in which fragments of the C. difficile toxin A repeat domain were expressed as genetic fusions with the C. botulinum C fragment gene using the pMAL-c expression vector (New England BioLabs). The pMAL-c expression vector generates fusion proteins which contain the MBP at the amino-terminal end of the protein. A construct, pMBot, in which the C. botulinum C fragment gene was expressed as a fusion with only the MBP was constructed (Figure 25). Fusion protein expression was induced from E. coli strains harboring the above plasmids, and induced protein was affinity purified on an amylose resin column.

i) Construction Of pBlueBot

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In order to facilitate the cloning of the *C. botulinum* C fragment gene sequences into a number of desired constructs, the botulinal gene sequences were removed from pAlterBot and were inserted into the pBluescript plasmid (Stratagene) to generate pBlueBot (Figure 25). pBlueBot was constructed as follows. Bacteria containing the pAlterBot plasmid were grown in medium containing tetracycline and plasmid DNA was isolated using the QIAprep-spin Plasmid Kit (Qiagen). One microgram of pAlterBot DNA was digested with *NcoI* and the resulting 3' recessed sticky end was made blunt using the Klenow fragment of DNA polymerase I (here after the Klenow fragment). The pAlterBot DNA was then digested with *HindIII* to

release the botulinal gene sequences (the Bot insert) as a blunt (filled NcoI site)HindIII fragment. pBluescript vector DNA was prepared by digesting 200 ng of
pBluescript DNA with SmaI and HindIII. The digestion products from both plasmids
were resolved on an agarose gel. The appropriate fragments were removed from the
gel, mixed and purified utilizing the Prep-a-Gene kit (BioRad). The eluted DNA was
then ligated using T4 DNA ligase and used to transform competent DH5α cells
(Gibco-BRL). Host cells were made competent for transformation using the calcium
chloride protocol of Sambrook et al., supra at 1.82-1.83. Recombinant clones were
isolated and confirmed by restriction digestion using standard recombinant molecular
biology techniques (Sambrook et al, supra). The resultant clone, pBlueBot, contains
several useful unique restriction sites flanking the Bot insert (i.e., the C. botulinum C
fragment sequences derived from pAlterBot) as shown in Figure 25.

ii) Construction Of C. difficile /C. botulinum / MBP Fusion Proteins

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Constructs encoding fusions between the C. difficile toxin A gene and the C. botulinum C fragment gene and the MBP were made utilizing the same recombinant DNA methodology outlined above; these fusion proteins contained varying amounts of the C. difficile toxin A repeat domain.

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The pMABot clone contains a 2.4 kb insert derived from the *C. difficile* toxin A gene fused to the Bot insert (*i.e.*, the *C. botulinum* C fragment sequences derived from pAlterBot). pMABot (Figure 25) was constructed by mixing gel-purified DNA from *Notl/Hind*III digested pBlueBot (the 1.2 kb Bot fragment), *Spel/Notl* digested pPA1100-2680 (the 2.4 kb *C. difficile* toxin A repeat fragment) and *Xbal/Hind*III digested pMAL-c vector. Recombinant clones were isolated, confirmed by restriction digestion and purified using the QIAprep-spin Plasmid Kit (Qiagen). This clone expresses the toxin A repeats and the botulinal C fragment protein sequences as an inframe fusion with the MBP.

The pMCABot construct contains a 1.0 kb insert derived from the *C. difficile* toxin A gene fused to the Bot insert (i.e, the *C. botulinum* C fragment sequences derived from pAlterBot). pMCABot was constructed by digesting the pMABot clone with *Eco*RI to remove the 5' end of the *C. difficile* toxin A repeat (see Figure 25, the pMAL-c vector contains a *Eco*RI site 5' to the *C. difficile* insert in the pMABot clone). The restriction sites were filled and religated together after gel purification. The resultant clone (pMCABot, Figure 25) generated an in-frame fusion between the MBP and the remaining 3' portion of the *C. difficile* toxin A repeat domain fused to the Bot gene.

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The pMNABot clone contains the 1 kb Spel/EcoRI (filled) fragment from the C. difficile toxin A repeat domain (derived from clone pPA1100-2680) and the 1.2 kb C. botulinum C fragment gene as a NcoI (filled)/HindIII fragment (derived from pAlterBot). These two fragments were inserted into the pMAL-c vector digested with Xbal/HindIII. The two insert fragments were generated by digestion of the appropriate plasmid with EcoRI (pPA1100-2680) or NcoI (pAlterBot) followed by treatment with the Klenow fragment. After treatment with the Klenow fragment, the plasmids were digested with the second enzyme (either SpeI or HindIII). All three fragments were gel purified, mixed and Prep-a-Gene purified prior to ligation. Following ligation and transformation, putative recombinants were analyzed by restriction analysis; the EcoRI site was found to be regenerated at the fusion junction, as was predicted for a fusion between the filled EcoRI and NcoI sites.

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A construct encoding a fusion protein between the botulinal C fragment gene and the MBP gene was constructed (i.e., this fusion lacks any C. difficile toxin A gene sequences) and termed pMBot. The pMBot construct was made by removal of the C. difficile toxin A sequences from the pMABot construct and fusing the C fragment gene sequences to the MBP. This was accomplished by digestion of pMABot DNA with StuI (located in the pMALc polylinker 5' to the XbaI site) and XbaI (located 3' to the NotI site at the toxA-Bot fusion junction), filling in the XbaI site using the Klenow fragment, gel purifying the desired restriction fragment, and ligating the blunt ends to

circularize the plasmid. Following ligation and transformation, putative recombinants were analyzed by restriction mapping of the Bot insert (i.e, the C. botulinum C fragment sequences).

b) Expression Of C. botulinum C Fragment Fusion Proteins In E. coli

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Large scale (1 liter) cultures of the pMAL-c vector, and each recombinant construct described above in (a) were grown, induced, and soluble protein fractions were isolated as described in Example 18. The soluble protein extracts were chromatographed on amylose affinity columns to isolate recombinant fusion protein. The purified recombinant fusion proteins were analyzed by running samples on SDS-PAGE gels followed by Coomassie staining and by Western blot analysis as described [Williams et al, (1994) supra]. In brief, extracts were prepared and chromatographed in column buffer (10 mM NaPO₄, 0.5 M NaCl, 10 mM β-mercaptoethanol, pH 7.2) over an amylose resin (New England Biolabs) column, and eluted with column buffer containing 10 mM maltose as described [Williams, et al. (1994), supra]. An SDS-PAGE gel containing the purified protein samples stained with Coomassie blue is shown in Figure 26.

In Figure 26, the following samples were loaded. Lanes 1-6 contain protein purified from *E. coli* containing the pMAL-c, pPA1870-2680, pMABot, pMNABot, pMCABot and pMBot plasmids, respectively. Lane 7 contains broad range molecular weight protein markers (BioRad).

The protein samples were prepared for electrophoresis by mixing 5 μl of eluted protein with 5 μl of 2X SDS-PAGE sample buffer (0.125 mM Tris-HCl, pH 6.8, 2 mM EDTA, 6% SDS, 20% glycerol, 0.025% bromophenol blue; β-mercaptoethanol is added to 5% before use). The samples were heated to 95°C for 5 min, then cooled and loaded on a 7.5% agarose SDS-PAGE gel. Broad range molecular weight protein markers were also loaded to allow estimation of the MW of identified fusion proteins.

After electrophoresis, protein was detected generally by staining the gel with Coomassie blue.

In all cases the yields were in excess of 20 mg fusion protein per liter culture (see Table 36) and, with the exception of the pMCABot protein, a high percentage (i.e., greater than 20-50% of total eluted protein) of the eluted fusion protein was of a MW predicted for the full length fusion protein (Figure 26). It was estimated (by visual inspection) that less than 10% of the pMCABot fusion protein was expressed as the full length fusion protein.

TABLE 36

Yield Of Affinity Purified C. botulinum C Fragment / MBP Fusion Proteins

Construct	Yield (mg/liter of Culture)	Percentage Of Total Soluble Protein
pMABot	. 24	5.0
pMCABot	34	5.0
pMNABot	40	5.5
pMBot	22	5.0
pMA1870-2680	40	4.8

These results demonstrate that high level expression of intact C. botulinum C fragment/C. difficile toxin A fusion proteins in E. coli is feasible using the pMAL-c expression system. These results are in contrast to those reported by H. F. LaPenotiere, et al. (1993), supra. In addition, these results show that it is not necessary to fuse the botulinal C fragment gene to the C. difficile toxin A gene in order to produce a soluble fusion protein using the pMAL-c system in E. coli.

In order to determine whether the above-described botulinal fusion proteins were recognized by anti-C. botulinum toxin A antibodies, Western blots were performed. Samples containing affinity-purified proteins from E. coli containing the pMABot, pMCABot, pMNABot, pMBot, pMA1870-2680 or pMALc plasmids were analyzed. SDS-PAGE gels (7.5% acrylamide) were loaded with protein samples purified from each expression construct. After electrophoresis, the gels were blotted

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and protein transfer was confirmed by Ponceau S staining (as described in Example 12b).

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Following protein transfer, the blots were blocked by incubation for 1 hr at 20°C in blocking buffer [PBST (PBS containing 0.1% Tween 20 and 5% dry milk)]. The blots were then incubated in 10 ml of a solution containing the primary antibody; this solution comprised a 1/500 dilution of an anti-C. botulinum toxin A IgY PEG prep (described in Example 3) in blocking buffer. The blots were incubated for 1 hr at room temperature in the presence of the primary antibody. The blots were washed and developed using a rabbit anti-chicken alkaline phosphatase conjugate (Boehringer Mannheim) as the secondary antibody as follows. The rabbit anti-chicken antibody was diluted to 1 µg/ml in blocking buffer (10 ml final volume per blot) and the blots were incubated at room temperature for 1 hour in the presence of the secondary antibody. The blots were then washed successively with PBST, BBS-Tween and 50 mM Na₂CO₃, pH 9.5. The blots were then developed in freshly-prepared alkaline phosphatase substrate buffer (100 µg/ml nitro blue tetrazolium, 50 µg/ml 5-bromo-chloro-indolylphosphate, 5 mM MgCl₂ in 50 mM Na₂CO₃, pH 9.5). Development was stopped by flooding the blots with distilled water and the blots were air dried.

This Western blot analysis detected anti-C. botulinum toxin reactive proteins in the pMABot, pMCABot, pMNABot and pMBot protein samples (corresponding to the predicted full length proteins identified above by Coomassie staining in Figure 26), but not in the pMA1100-2680 or pMALc protein samples.

These results demonstrate that the relevant fusion proteins purified on an amylose resin as described above in section a) contained immunoreactive C botulinum C fragment protein as predicted.

EXAMPLE 23

Generation Of Neutralizing Antibodies By Nasal Administration Of pMBot Protein

The ability of the recombinant botulinal toxin proteins produced in Example 22 to stimulate a systemic immune response against botulinal toxin epitopes was assessed. This example involved: a) the evaluation of the induction of serum IgG titers produced by nasal or oral administration of botulinal toxin-containing *C. difficile* toxin A fusion proteins and b) the *in vivo* neutralization of *C. botulinum* type A neurotoxin by anti- recombinant *C. botulinum* C fragment antibodies.

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a) Evaluation Of The Induction Of Serum IgG Titers

Produced By Nasal Or Oral Administration Of

Botulinal Toxin-Containing C. difficile Toxin A Fusion

Proteins

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Six groups containing five 6 week old CF female rats (Charles River) per group were immunized nasally or orally with one of the following three combinations using protein prepared in Example 22: (1) 250 µg pMBot protein per rat (nasal and oral); 2) 250 µg pMABot protein per rat (nasal and oral); 3) 125 µg pMBot admixed with 125 µg pMA1870-2680 per rat (nasal and oral). A second set of 5 groups containing 3 CF female rats/group were immunized nasally or orally with one of the following combinations (4) 250 µg pMNABot protein per rat (nasal and oral) or 5) 250 µg pMAL-c protein per rat (nasal and oral).

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The fusion proteins were prepared for immunization as follows. The proteins (in column buffer containing 10 mM maltose) were diluted in 0.1 M carbonate buffer, pH 9.5 and administered orally or nasally in a 200 µl volume. The rats were lightly sedated with ether prior to administration. The oral dosing was accomplished using a 20 gauge feeding needle. The nasal dosing was performed using a P-200 micropipettor (Gilson). The rats were boosted 14 days after the primary immunization using the techniques described above and were bled 7 days later. Rats from each group were

lightly etherized and bled from the tail. The blood was allowed to clot at 37°C for 1 hr and the serum was collected.

The serum from individual rats was analyzed using an ELISA to determine the anti-C. botulinum type A toxin IgG serum titer. The ELISA protocol used is a modification of that described in Example 13c. Briefly, 96-well microtiter plates (Falcon, Pro-Bind Assay Plates) were coated with C. botulinum type A toxoid (prepared as described in Example 3a) by placing 100 µl volumes of C. botulinum type A toxoid at 2.5 µg/ml in PBS containing 0.005% thimerosal in each well and incubating overnight at 4°C. The next morning, the coating suspensions were decanted and all wells were washed three times using PBS.

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In order to block non-specific binding sites, 100 µl of blocking solution [0.5%] BSA in PBS] was then added to each well and the plates were incubated for 1 hr at 37°C. The blocking solution was decanted and duplicate samples of 150 µl of diluted rat serum added to the first well of a dilution series. The initial testing serum dilution was 1:30 in blocking solution containing 0.5% Tween 20 followed by 5-fold dilutions into this solution. This was accomplished by serially transferring 30 µl aliquots to 120 ul blocking solution containing 0.5% Tween 20, mixing, and repeating the dilution into a fresh well. After the final dilution, 30 µl was removed from the well such that all wells contained 120 µl final volume. A total of 3 such dilutions were performed (4 wells total). The plates were incubated 1 hr at 37°C. Following this incubation, the serially diluted samples were decanted and the wells were washed six times using PBS containing 0.5% Tween 20 (PBST). To each well, 100 µl of a rabbit anti-Rat IgG alkaline phosphatase (Sigma) diluted (1/1000) in blocking buffer containing 0.5% Tween 20 was added and the plate was incubated for 1 hr at 37°C. The conjugate solutions were decanted and the plates were washed as described above, substituting 50 mM Na₂CO₃, pH 9.5 for the PBST in the final wash. The plates were developed by the addition of 100 µl of a solution containing 1 mg/ml para-nitro phenyl phosphate (Sigma) dissolved in 50 mM Na₂CO₃, 10 mM MgCl₂, pH 9.5 to each well, and incubating the plates at room temperature in the dark for 5-45 min. The absorbency of each well was measured at 410 nm using a Dynatech MR 700 plate reader. The

results are summarized in Tables 37 and 38 and represent mean serum reactivities of individual mice.

TABLE 37

Determination Of Anti-C. botulinum Type A Toxin Serum IgG Titers
Following Immunization With C. botulinum C Fragment-Containing Fusion Proteins

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Route of Immunization		Nasal Oral					
İmmunogen	PRE- IMMUNE	pMBat	pMBot & pMA1870- 2680	рМАВоι	pMBot	pMBot& pMA1870- 2680	рМАВоі
Dilution							
1:30	0.080	1.040	1.030	0.060	0.190	0.080	0.120
1:150	0.017	0.580	0.540	0.022	0.070	0.020	0.027
1:750	0.009	0.280	0.260	0.010	0.020	0.010	0.014
l:3750	0.007	0.084	0.090	0.009	0.009	0.010	0.007
# Rats Tested		5	5	· 5	5	2	2

 Numbers represent the average values obtained from two ELISA plates, standardized utilizing the preimmune control.

TABLE 38

Determination Of Anti-C. botulinum Type A Toxin Serum IgG Titers
Following Immunization With C. botulinum C Fragment-Containing Fusion Proteins

Route of Immunization		Nasal		Oral	
Immunogen	PRE; IMMUNE	pMBot	pMABot	pMNABot	pMNABot
Dilution					
1:30	0.040	0.557	0.010	0.015	0.010
1:150	0.009	0.383	0.001	0.003	0.002
1:750	0.001	0.140	0.000	0.000	0.000
1:3750	0.000	0.040	0.000	0.000	0.000
# Rats Tested		1	ì	3	3

The above ELISA results demonstrate that reactivity against the botulinal fusion proteins was strongest when the route of administration was nasal; only weak responses were stimulated when the botulinal fusion proteins were given orally.

Nasally delivered pMbot and pMBot admixed with pMA1870-2680 invoked the greatest serum IgG response. These results show that only the pMBot protein is necessary to induce this response, since the addition of the pMA1870-2680 protein did not enhance antibody response (Table 37). Placement of the C. difficile toxin A fragment between the MBP and the C. botulinum C fragment protein dramatically reduced anti-bot IgG titer (see results using pMABot, pMCABot and pMNABot proteins).

This study demonstrates that the pMBot protein induces a strong serum IgG response directed against C. botulinum type A toxin when nasally administered.

b) In Vivo Neutralization Of C. botulinum Type A Neurotoxin By Anti- Recombinant C. botulinum C Fragment Antibodies

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The ability of the anti-C. botulinum type: A toxin antibodies generated by nasal administration of recombinant botulinal fusion proteins in rats (Example 22) to neutralize C. botulinum type A toxin was tested in a mouse neutralization model. The mouse model is the art accepted method for detection of botulinal toxins in body fluids and for the evaluation of anti-botulinal antibodies [E.J. Schantz and D.A. Kautter, J. Assoc. Off. Anal. Chem. 61:96 (1990) and Investigational New Drug (BB-IND-3703) application by the Surgeon General of the Department of the Army to the Federal Food and Drug Administration]. The anti-C. botulinum type A toxin antibodies were prepared as follows.

Rats from the group given pMBot protein by nasal administration were boosted a second time with 250 µg pMBot protein per rat and serum was collected 7 days later. Serum from one rat from this group and from a preimmune rat was tested for anti-C. botulinum type A toxin neutralizing activity in the mouse neutralization model described below.

The LD₅₀ of a solution of purified *C. botulinum* type A toxin complex, obtained from Dr. Eric Johnson (University of Wisconsin Madison), was determined using the

intraperitoneal (IP) method of Schantz and Kautter [J. Assoc. Off. Anal. Chem. 61:96 (1978)] using 18-22 gram female ICR mice and was found to be 3500 LD₅₀/ml. The determination of the LD₅₀ was performed as follows. A Type A toxin standard was prepared by dissolving purified type A toxin complex in 25 mM sodium phosphate buffer, pH 6.8 to yield a stock toxin solution of 3.15 x 10⁷ LD₅₀/mg. The OD₂₇₈ of the solution was determined and the concentration was adjusted to 10-20 μg/ml. The toxin solution was then diluted 1:100 in gel-phosphate (30 mM phosphate, pH 6.4; 0.2% gelatin). Further dilutions of the toxin solution were made as shown below in Table 39. Two mice were injected IP with 0.5 ml of each dilution shown and the mice were observed for symptoms of botulism for a period of 72 hours.

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TABLE 39

Determination Of The LD₅₀ Of Purified C. botulinum Type A Toxin Complex

Dilution	Number Dead At 72 hr					
1:320	2/2					
1:640	2/2					
1:1280	2/2					
1:2560	0/2 (sick after 72 hr)					
1:5120	0/2 (no symptoms)					

From the results shown in Table 39, the toxin titer was assumed to be between $2560 \text{ LD}_{50}/\text{ml}$ and $5120 \text{ LD}_{50}/\text{ml}$ (or about $3840 \text{ LD}_{50}/\text{ml}$). This value was rounded to $3500 \text{ LD}_{50}/\text{ml}$ for the sake of calculation.

The amount of neutralizing antibodies present in the serum of rats immunized nasally with pMBot protein was then determined. Serum from two rats boosted with pMBot protein as described above and preimmune serum from one rat was tested as follows. The toxin standard was diluted 1:100 in gel-phosphate to a final concentration of 350 LD₅₀/ml. One milliliter of the diluted toxin standard was mixed with 25 µl of serum from each of the three rats and 0.2 ml of gel-phosphate. The mixtures were incubated at room temperature for 30 min with occasional mixing. Each of two mice were injected with IP with 0.5 ml of the mixtures. The mice were

observed for signs of botulism for 72 hr. Mice receiving serum from rats immunized with pMBot protein neutralized this challenge dose. Mice receiving preimmune rat serum died in less than 24 hr.

The amount of neutralizing anti-toxin antibodies present in the serum of rats immunized with pMBot protein was then quantitated. Serum antibody titrations were performed by mixing 0.1 ml of each of the antibody dilutions (see Table 40) with 0.1 ml of a 1:10 dilution of stock toxin solution (3.5 x 10⁴ LD₅₀/ml) with 1.0 ml of gel-phosphate and injecting 0.5 ml IP into 2 mice per dilution. The mice were then observed for signs of botulism for 3 days (72 hr). The results are tabulated in Table 39.

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As shown in Table 40 pMBot serum neutralized *C. botulinum* type A toxin complex when used at a dilution of 1:320 or less. A mean neutralizing value of 168 IU/ml was obtained for the pMBot serum (an IU is defined as 10,000 mouse LD₅₀). This value translates to a circulating serum titer of about 3.7 IU/mg of serum protein. This neutralizing titer is comparable to the commercially available bottled concentrated (Connaught Laboratories, Ltd.) horse anti-*C. botulinum* antiserum. A 10 ml vial of Connaught antiserum contains about 200 mg/ml of protein; each ml can neutralize 750 IU of *C. botulinum* type A toxin. After administration of one vial to a human, the circulating serum titer of the Connaught preparation would be approximately 25 IU/ml assuming an average serum volume of 3 liters). Thus, the circulating anti-*C. botulinum* titer seen in rats nasally immunized with pMBot protein (168 IU/ml) is 6.7 time higher than the necessary circulation titer of anti-*C. botulinum* antibody needed to be protective in humans.

TABLE 40
Quantitation Of Neutralizing Antibodies In pMBot Sera

P.	pMBot*					
Dilution	Rat I	Rat 2				
1:20	2/2	2/2				
1:40	2/2	2/2				
1:80	2/2	2/2				
1:160	. 2/2	2/2				
1:320	2/2 ^b	2/2 ⁶				
1:640	0/2	0/2				
1:1280	0/2	0/2				
1:2560	0/2	0/2				

- Numbers represent the number of mice surviving at 72 hours which received serum taken from rats immunized with the pMBot protein.
- These mice survived but were sick after 72 hr.

These results demonstrate that antibodies capable of neutralizing *C. botulinum* type A toxin are induced when recombinant *C. botulinum* C fragment fusion protein produced in *E. coli* is used as an immunogen.

EXAMPLE 24

Production Of Soluble C. botulinum C Fragment
Protein Substantially Free Of Endotoxin Contamination

Example 23 demonstrated that neutralizing antibodies are generated by immunization with the pMBot protein expressed in *E. coli*. These results showed that the pMBot fusion protein is a good vaccine candidate. However, immunogens suitable for use as vaccines should be pyrogen-free in addition to having the capability of inducing neutralizing antibodies. Expression clones and conditions that facilitate the production of *C. botulinum* C fragment protein for utililization as a vaccine were developed.

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The example involved: (a) determination of pyrogen content of the pMBot protein; (b) generation of *C. botulinum* C fragment protein free of the MBP; (c) expression of *C. botulinum* C fragment protein using various expression vectors; and (d) purification of soluble *C. botulinum* C fragment protein substantially free of significant endotoxin contamination.

a) Determination Of The Pyrogen Content Of The pMBot Protein

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In order to use a recombinant antigen as a vaccine in humans or other animals, the antigen preparation must be shown to be free of pyrogens. The most significant pyrogen present in preparations of recombinant proteins produced in gram-negative bacteria, such as *E. coli*, is endotoxin [F.C. Pearson, *Pyrogens: endotoxins, LAL testing and depyrogentaion*, (1985) Marcel Dekker, New York, pp. 23-56]. To evaluate the utility of the pMBot protein as a vaccine candidate, the endotoxin content in MBP fusion proteins was determined.

The endotoxin content of recombinant protein samples was assayed utilizing the Limulus assay (LAL kit; Associates of Cape Cod) according to the manufacturer's instructions. Samples of affinity-purified pMal-c protein and pMA1870-2680 were found to contain high levels of endotoxin [>50,000 EU/mg protein; EU (endotoxin unit)]. This suggested that MBP- or toxin A repeat-containing fusions with the botulinal C fragment should also contain high levels of endotoxin. Accordingly, removal of endotoxin from affinity-purified pMal-c and pMBot protein preparations was attempted as follows.

Samples of pMal-c and pMBot protein were depyrogenated with polymyxin to determine if the endotoxin could be easily removed. The following amount of protein was treated: 29 ml at 4.8 OD₂₈₀/ml for pMal-c and 19 mls at 1.44 OD₂₈₀/ml for pMBot. The protein samples were dialyzed extensively against PBS and mixed in a 50 ml tube (Falcon) with 0.5 ml PBS-equilibrated polymyxin B (Affi-Prep Polymyxin, BioRad). The samples were allowed to mix by rotating the tubes overnight at 4°C.

The polymyxin was pelleted by centrifugation for 30 min in a bench top centrifuge at maximum speed (approximately 2000 x g) and the supernatant was removed. The recovered protein (in the supernatant) was quantified by OD₂₈₀, and the endotoxin activity was assayed by LAL. In both cases only approximately 1/3 of the input protein was recovered and the polymyxin-treated protein retained significant endotoxin contamination (approximately 7000 EU/mg of pMBot).

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The depyrogenation experiment was repeated using an independently purified pMal-c protein preparation and similar results were obtained. From these studies it was concluded that significant levels of endotoxin copurifies with these MBP fusion proteins using the amylose resin. Furthermore, this endotoxin cannot be easily removed by polymyxin treatment.

These results suggest that the presence of the MBP sequences on the fusion protein complicated the removal of endotoxin from preparations of the pMBot protein.

b) Generation Of C. botulinum C Fragment Protein Free Of The MBP

It was demonstrated that the pMBot fusion protein could not be easily purified from contaminating endotoxin in section a) above. The ability to produce a pyrogen-free (e.g., endotoxin-free) preparation of soluble botulinal C fragment protein free of the MBP tag was next investigated. The pMBot expression construct was designed to facilitate purification of the botulinal C fragment from the MBP tag by cleavage of the fusion protein by utilizing an engineered Factor Xa cleavage site present between the MBP and the botulinal C fragment. The Factor Xa cleavage was performed as follows.

Factor Xa (New England Biolabs) was added to the pMBot protein (using a 0.1-1.0% Factor Xa/pMBot protein ratio) in a variety of buffer conditions [e.g., PBS-NaCl (PBS containing 0.5 M NaCl), PBS-NaCl containing 0.2% Tween 20, PBS, PBS containing 0.2% Tween 20, PBS-C (PBS containing 2 mM CaCl₂), PBS-C containing either 0.1 or 0.5 % Tween 20, PBS-C containing either 0.1 or 0.5% NP-40, PBS-C containing either 0.1 or 0.5% Triton X-100, PBS-C containing 0.1% sodium

deoxycholate, PBS-C containing 0.1% SDS]. The Factor Xa digestions were incubated for 12-72 hrs at room temperature.

The extent of cleavage was assessed by Western blot or Coornassie blue staining of proteins following electrophoresis on denaturing SDS-PAGE gels, as described in Example 22. Cleavage reactions (and control samples of uncleaved pMBot protein) were centrifuged for 2 min in a microfuge to remove insoluble protein prior to loading the samples on the gel. The Factor Xa treated samples were compared with uncleaved, uncentrifuged pMBot samples on the same gel. The results of this analysis is summarized below.

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1) Most (about 90%) pMBot protein could be removed by centrifugation, even when uncleaved control samples were utilized. This indicated that the pMBot fusion protein was not fully soluble (i.e., it exists as a suspension rather than as a solution). [This result was consistent with the observation that most affinity-purified pMBot protein precipitates after long term storage (>2 weeks) at 4°C. Additionally, the majority (i.e., 75%) of induced pMBot protein remains in the pellet after sonication and clarification of the induced *E. coli*. Resuspension of these insoluble pellets in PBS followed by sonication results in partial solubilization of the insoluble pMBot protein in the pellets.]

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2) The portion of pMBot protein that is fully in solution (about 10% of pMBot protein) is completely cleaved by Factor Xa, but the cleaved (released) botulinal C fragment is relatively insoluble such that only the cleaved MBP remains fully in solution.

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reducing effective cleavage. Conditions that effectively solubilized the cleaved botulinal C fragment were not identified.

4) The use of 0.1% SDS in the buffer used for Factor Xa cleavage

None of the above reaction conditions enhanced solubility without also

enhanced the solubility of the pMBot protein (all of pMBot protein was soluble).

However, the presence of the SDS prevented any cleavage of the fusion protein with Factor Xa.

5) Analysis of pelleted protein from the cleavage reactions indicated that both full length pMBot (i.e., uncleaved) and cleaved botulinal C fragment protein precipitated during incubation.

These results demonstrate that purification of soluble botulinal C fragment protein after cleavage of the pMBot fusion protein is complicated by the insolubility of both the pMBot protein and the cleaved botulinal C fragment protein.

c) Expression Of C. botulinum C Fragment Using Various Expression Vectors

In order to determine if the solubility of the botulinal C fragment was enhanced by expressing the C fragment protein as a native protein, an N-terminal His-tagged protein or as a fusion with glutathione-S-transferase (GST), alternative expression plasmids were constructed. These expression constructs were generated utilizing the methodologies described in Example 22. Figure 27 provides a schematic representation of the vectors described below.

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In Figure 27, the following abbreviations are used. pP refers to the pET23 vector. pHIS refers to the pETHisa vector. pBlue refers to the pBluescript vector. pM refers to the pMAL-c vector and pG refers to the pGEX3T vector (described in Example 11). The solid black lines represent *C. botulinum* C fragment gene sequences; the solid black ovals represent the MBP; the hatched ovals represent GST; "HHHHH" represents the poly-histidine tag. In Figure 27, when the name for a restriction enzyme appears inside parenthesis, this indicates that the restriction site was destroyed during construction. An asterisk appearing with the name for a restriction enzyme indicates that this restriction site was recreated at a cloning junction.

i) Construction Of pPBot

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In order to express the *C. botulinum* C fragment as a native (i.e., non-fused) protein, the pPBot plasmid (shown schematically in Figure 27) was constructed as follows. The C fragment sequences present in pAlterBot (Example 22) were removed by digestion of pAlterBot with *Ncol* and *HindIII*. The *Ncol/HindIII* C fragment insert

was ligated to pETHisa vector (described in Example 18b) which was digested with Ncol and HindIII. This ligation creates an expression construct in which the Ncolencoded methionine of the botulinal C fragment is the initiator codon and directs expression of the native botulinal C fragment. The ligation products were used to transform competent BL21(DE3)pLysS cells (Novagen). Recombinant clones were identified by restriction mapping.

ii) Construction Of pHisBot

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In order to express the *C. botulinum* C fragment containing a poly-histidine tag at the amino-terminus of the recombinant protein, the pHisBot plasmid (shown schematically in Figure 27) was constructed as follows. The *Ncol/Hind*III botulinal C fragment insert from pAlterbot was ligated into the pETHisa vector which was digested with *NheI* and *Hind*III. The *NcoI* (on the C fragment insert) and *NheI* (on the pETHisa vector) sites were filled in using the Klenow fragment prior to ligation; these sites were then blunt end ligated (the *NdeI* site was regenerated at the clone junction as predicted). The ligation products were used to transform competent BL21(DE3)pLysS cells and recombinant clones were identified by restriction mapping.

The resulting pHisBot clone expresses the botulinal C fragment protein with a histidine-tagged N-terminal extension having the following sequence: MetGlyHisHis HisHisHisHisHisHisHisHisSerSerGlyHisIleGluGlyArgHisMetAla, (SEQ ID NO:24); the amino acids encoded by the botulinal C fragment gene are underlined and the vector encoded amino acids are presented in plain type. The nucleotide sequence present in the pETHisa vector which encodes the pHisBot fusion protein is listed in SEQ ID NO:25. The amino acid sequence of the pHisBot protein is listed in SEQ ID NO:26.

iii) Construction Of pGBot

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The botulinal C fragment protein was expressed as a fusion with the glutathione-S-transferase protein by constructing the pGBot plasmid (shown schematically in Figure 27). This expression construct was created by cloning the NotI/Sall C fragment insert present in pBlueBot (Example 22) into the pGEX3T vector which was digested with Smal and Xhol. The NotI site (present on the botulinal fragment) was made blunt prior to ligation using the Klenow fragment. The ligation products were used to transform competent BL21 cells.

Each of the above expression constructs were tested by restriction digestion to confirm the integrity of the constructs.

Large scale (1 liter) cultures of pPBot [BL21(DE3)pLysS host], pHisBot [BL21(DE3)pLysS host] and pGBot (BL21 host) were grown in 2X YT medium and induced (using IPTG to 0.8-1.0 mM) for 3 hrs as described in Example 22. Total, soluble and insoluble protein preparations were prepared from 1 ml aliquots of each large scale culture [Williams et al. (1994), supra] and analyzed by SDS-PAGE. No obvious induced band was detectable in the pPBot or pHisBot samples by Coomassie staining, while a prominent insoluble band of the anticipated MW was detected in the pGBot sample. Soluble lysates of the pGBot large scale (resuspended in PBS) or pHisBot large scale [resuspended in Novagen 1X binding buffer (5 mM imidazole, 0.5 M NaCl, 20 mM Tris-HCl, pH 7.9)] cultures were prepared and used to affinity purify soluble affinity-tagged protein as follows.

The pGBot lysate was affinity purified on a glutathione-agarose resin (Pharmacia) exactly as described in Smith and Corcoran [Current Protocols in Molecular Biology, Supplement 28 (1994), pp. 16.7.1-16.7.7]. The pHisBot protein was purified on the His-Bind resin (Novagen) utilizing the His-bind buffer kit (Novagen) exactly as described by manufacturer.

Samples from the purification of both the pGBot and pHisBot proteins (including uninduced, induced, total, soluble, and affinity-purified eluted protein) were resolved on SDS-PAGE gels. Following electrophoresis, proteins were analyzed by

Coomassie staining or by Western blot detection utilizing a chicken anti-C. botulinum Type A toxoid antibody (as described in Example 22).

These studies showed that the pGBot protein was almost entirely insoluble under the utilized conditions, while the pHisBot protein was soluble. Affinity purification of the pHisBot protein on this first attempt was inefficient, both in terms of yield (most of the immunoreactive botulinal protein did not bind to the His-bind resin) and purity (the botulinal protein was estimated to comprise approximately 20% of the total eluted protein).

d) Purification Of Soluble C. botulinum C Fragment Protein Substantially Free Of Endotoxin Contamination

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The above studies showed that the pHisBot protein was expressed in $E.\ coli$ as a soluble protein. However, the affinity purification of this protein on the His-bind resin was very inefficient. In order to improve the affinity purification of the soluble pHisBot protein (in terms of both yield and purity), an alternative poly-histidine binding affinity resin (Ni-NTA resin; Qiagen) was utilized. The Ni-NTA resin was reported to have a superior binding affinity ($K_d=1 \times 10^{-13}$ at pH 8.0; Qiagen user manual) relative to the His-bind resin.

A soluble lysate (in Novagen 1X binding buffer) from an induced 1 liter 2X YT culture was prepared as described above. Briefly, the culture of pHisBot [Bl21(DE3)pLysS host] was grown at 37°C to an OD₆₀₀ of 0.7 in 1 liter of 2X YT medium containing 100 μg/ml ampicillin, 34 μg/ml chloramphenicol and 0.2% glucose. Protein expression was induced by the addition of IPTG to 1 mM. Three hours after the addition of the IPTG, the cells were cooled for 15 min in a ice water bath and then centrifuged 10 min at 5000 rpm in a JA10 rotor (Beckman) at 4°C. The pellets were resuspended in a total volume of 40 mls Novagen 1X binding buffer (5 mM imidazole, 0.5 M NaCl, 20 mM Tris-HCl, pH 7.9), transferred to two 35 ml Oakridge tubes and frozen at -70°C for at least 1 hr. The tubes were thawed and the cells were lysed by sonication (4 X 20 second bursts using a Branson Sonifier 450

with a power setting of 6-7) on ice. The suspension was clarified by centrifugation for 20 min at 9,000 rpm (10,000 x g) in a JA-17 rotor (Beckman).

The soluble lysate was brought to 0.1% NP40 and then was batch absorbed to 7 ml of a 1:1 slurry of Ni-NTA resin:binding buffer by stirring for 1 hr at 4°C. The slurry was poured into a column having an internal diameter of 1 or 2.5 cm (BioRad). The column was then washed sequentially with 15 mls of Novagen 1X binding buffer containing 0.1% NP40, 15 ml of Novagen 1X binding buffer, 15 ml wash buffer (60 mM imidazole, 0.5 M NaCl, 20 mM Tris-HCl, pH 7.9) and 15 ml NaHPO₄ wash buffer (50 mM NaHPO₄, pH 7.0, 0.3 M NaCl, 10 % glycerol). The bound protein was eluted by protonation of the resin using elution buffer (50 mM NaHPO₄, pH 4.0, 0.3 M NaCl, 10 % glycerol). The eluted protein was stored at 4°C.

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Samples of total, soluble and eluted protein were resolved by SDS-PAGE. Protein samples were prepared for electrophoresis as described in Example 22b. Duplicate gels were stained with Coomassie blue to visualize the resolved proteins and C. botulinum type A toxin-reactive protein was detected by Western blot analysis as described in Example 22b. A representative Coomassie stained gel is shown in Figure 28. In Figure 28, the following samples were loaded on the 12.5% acrylamide gel. Lanes 1-4 contain respectively total protein, soluble protein, soluble protein present in the flow-through of the Ni-NTA column and affinity-purified pHisBot protein (i.e., protein released from the Ni-NTA resin by protonation). Lane 5 contains high molecular weight protein markers (BioRad).

The purification of pHisBot protein resulted in a yield of 7 mg of affinity purified protein from a 1 liter starting culture of BL21(DE3)pLysS cells harboring the pHisBot plasmid. The yield of purified pHisBot protein represented approximately 0.4% of the total soluble protein in the induced culture. Analysis of the purified pHisBot protein by SDS-PAGE revealed that at least 90-95% of the protein was present as a single band (Figure 28) of the predicted MW (50 kD). This 50 kD protein band was immunoreactive with anti-C. botulinum type A toxin antibodies. The extinction coefficient of the protein preparation was determined to be 1.4 (using the Pierce BCA assay) or 1.45 (using the Lowry assay) OD₂₈₀ per 1 mg/ml solution.

Samples of pH neutralized eluted pHisBot protein were resolved on a KB 803 HPLC column (Shodex). Although His-tagged proteins are retained by this sizing column (perhaps due to the inherent metal binding ability of the proteins), the relative mobility of the pHisBot protein was consistent with that expected for a non-aggregated protein in solution. Most of the induced pHisBot protein was determined to be soluble under the growth and solubilization conditions utilized above (i.e., greater than 90% of the pHisBot protein was found to be soluble as judged by comparison of the levels of pHisBot protein seen in total and soluble protein samples prepared from BL21(DE3)pLysS cells containing the pHisBot plasmid). SDS-PAGE analysis of samples obtained after centrifugation, extended storage at -20°C, and at least 2 cycles of freezing and thawing detected no protein loss (due to precipitation), indicating that the pHisBot protein is soluble in the elution buffer (i.e., 50 mM NaHPO4, pH 4.0, 0.3 M NaCl, 10 % glycerol).

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Determination of endotoxin contamination in the affinity purified pHisBot preparation (after pH neutralization) using the LAL assay (Associates of Cape Cod) detected no significant endotoxin contamination. The assay was performed using the endpoint chromogenic method (without diazo-coupling) according to the manufacturer's instructions. This method can detect concentrations of endotoxin greater than or equal to 0.03 EU/ml (EU refers to endotoxin units). The LAL assay was run using 0.5 ml of a solution comprising 0.5 mg pHisBot protein in 50 mM NaHPO₄, pH 7.0, 0.3 M NaCl, 10 % glycerol; 30-60 EU were detected in the 0.5 ml sample. Therefore, the affinity purified pHisBot preparation contains 60-120 EU/mg of protein. FDA Guidelines for the administration of parenteral drugs require that a composition to be administered to a human contain less than 5 EU/kg body weight (The average human body weight is 70 kg; therefore up to 349 EU units can be delivered in a parental dose.). Because very small amount of protein are administered in a vaccine preparation (generally in the range of 10-500 µg of protein), administration of affinity purified pHisBot containing 60-120 EU/mg protein would result in delivery of only a small percentage of the permissible endotoxin load. For example, administration of 10-500 µg of purified pHisBot to a 70 kg human, where

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the protein preparation contains 60 EU/mg protein, results in the introduction of only 0.6 to 30 EU [i.e., 0.2 to 8.6% of the maximum allowable endotoxin burden per parenteral dose (less than 5 EU/kg body weight)].

The above results demonstrate that endotoxin (LPS) does not copurify with the pHisBot protein using the above purification scheme. Preparations of recombinantly produced pHisBot protein containing lower levels of endotoxin (less than or equal to 2 EU/ mg recombinant protein) may be produced by washing the Ni-NTA column with wash buffer until the OD₂₈₀ returns to baseline levels (i.e., until no more UV-absorbing material comes off of the column).

The above results illustrate a method for the production and purification of soluble, botulinal C fragment protein substantially free of endotoxin.

EXAMPLE 25

Optimization Of The Expression And Purification Of pHisBot Protein

The results shown in Example 24d demonstrated that the pHisBot protein is an excellent candidate for use as a vaccine as it could be produced as a soluble protein in *E. coli* and could be purified free of pyrogen activity. In order to optimize the expression and purification of the pHisBot protein, a variety of growth and purification conditions were tested.

a) Growth Parameters

i) Host Strains

The influence of the host strain utilized upon the production of soluble pHisBot protein was investigated. A large scale purification of pHisBot was performed [as described in Example 24d above] using the BL21(DE3) host (Novagen) rather than the BL21(DE3)pLysS host. The deletion of the pLysS plasmid in the BL21(DE3) host yielded higher levels of expression due to de-repression of the plasmid's T7-lac promoter. However, the yield of affinity-purified soluble recombinant protein was very low (approximately 600 µg/ liter culture) when purified under conditions identical

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to those described in Example 24d above. This result was due to the fact that expression in the BL21(DE3) host yielded very high level expression of the pHisBot protein as insoluble inclusion bodies as shown by SDS-PAGE analysis of protein prepared from induced BL21(DE3) cultures (Figure 29, lanes 1-7, described below). These results demonstrate that the pHisBot protein is not inherently toxic to *E. coli* cells and can be expressed to high levels using the appropriate promoter/host combination.

Figure 29 shows a Coomassie blue stained SDS-PAGE gel (12.5% acrylamide) onto which extracts prepared from BL21(DE3) cells containing the pHisBot plasmid were loaded. Each lane was loaded with 2.5 μl protein sample mixed with 2.5 μl of 2X SDS sample buffer. The samples were handled as described in Example 22b. The following samples were applied to the gel. Lanes 1-7 contain protein isolated from the BL21(DE3) host. Lanes 8-14 contain proteins isolated from the BL21(DE3)pLysS host. Total protein was loaded in lanes 1, 2, 4, 6, 8, 10 and 12. Soluble protein was loaded in Lanes 3, 5, 7, 9, 11 and 13. Lane 1 contains protein from uninduced host cells. Lanes 2-13 contain protein from host cells induced for 3 hours. IPTG was added to a final concentration of 0.1 mM (Lanes 6-7), 0.3 mM (Lanes 4-5) or 1.0 mM (Lanes 2, 3, 8-13). The cultures were grown in LB broth (Lanes 8-9), 2X YT broth (Lanes 10-11) or terrific broth (Lanes 1-7, 12-13). The pHisBot protein seen in Lanes 3, 5 and 7 is insoluble protein which spilled over from Lanes 2, 4 and 6, respectively. High molecular weight protein markers (BioRad) were loaded in Lane 14.

A variety of expression conditions were tested to determine if the BL21(DE3) host could be utilized to express soluble pHisBot protein at suitably high levels (i.e., about 10 mg/ml). The conditions altered were temperature (growth at 37 or 30°C), culture medium (2X YT, LB or Terrific broth) and inducer levels (0.1, 0.3 or 1.0 mM IPTG). All combinations of these variables were tested and the induction levels and solubility was then assessed by SDS-PAGE analysis of total and soluble extracts [prepared from 1 ml samples as described in Williams et al., (1994), supra].

All cultures were grown in 15 ml tubes (Falcon #2057). All culture medium was prewarmed overnight at the appropriate temperature and were supplemented with

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100 μg/ml ampicillin and 0.2% glucose. Terrific broth contains 12 g/l bacto-tryptone, 24 g/l bacto-yeast extract and 100 ml/l of a solution comprising 0.17 M KH₂PO₄, 0.72 M K₂HPO₄. Cultures were grown in a incubator on a rotating wheel (to ensure aeration) to an OD₆₀₀ of approximately 0.4, and induced by the addition of IPTG. In all cases, high level expression of insoluble pHisBot protein was observed, regardless of temperature, medium or inducer concentration.

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The effect of varying the concentration of IPTG upon 2X YT cultures grown at 23°C was then investigated. IPTG was added to a final concentration of either 1 mM, 0.1 mM, 0.05 mM or 0.01 mM. At this temperature, similar levels of pHis Bot protein was induced in the presence of either 1 or 0.1 mM IPTG; these levels of expression was lower than that observed at higher temperatures. Induced protein levels were reduced at 0.05 mM IPTG and absent at 0.01 mM IPTG (relative to 1.0 and 0.1 mM IPTG inductions at 23°C). However, no conditions were observed in which the induced pHisBot protein was soluble in this host. Thus, although expression levels are superior in the BL21(DE3) host (as compared to the BL21(DE3)pLysS host), conditions that facilitate the production of soluble protein in this host could not be identified.

These results demonstrate that production of soluble pHisBot protein was achieved using the BL21(DE3)pLysS host in conjunction with the T7-lac promoter.

ii) Effect Of Varying Temperature,Medium And IPTG Concentration AndLength Of Induction

The effect growing the host cells in various mediums upon the expression of recombinant botulinal protein from the pHisBot expression construct [in the BL21(DE3)pLysS host] was investigated. BL21(DE3)pLysS cells containing the pHisBot plasmid were grown in either LB, 2X YT or Terrific broth at 37°C. The cells were induced using 1 mM IPTG for a 3 hr induction period. Expression of pHisBot protein was found to be the highest when the cells were grown in 2X YT broth (see Figure 29, lanes 8-13).

The cells were then grown at 30°C in 2X YT broth and the concentration of IPTG was varied from 1.0, 0.3 or 0.1 mM and the length of induction was either 3 or 5 hours. Expression of pHisBot protein was similar at all 3 inducer concentrations utilized and the levels of induced protein were higher after a 5 hr induction as compared to a 3 hr induction.

Using the conditions found to be optimal for the expression of pHisBot protein, a large scale culture was grown in order to provide sufficient material for a large scale purification of the pHisBot protein. Three 1 liter cultures were grown in 2X YT medium containing 100 µg/ml ampicillin, 34 µg/ml chloramphenicol and 0.2% glucose. The cultures were grown at 30°C and were induced with 1.0 mM IPTG for a 5 hr period. The cultures were harvested and a soluble lysate were prepared as described in Example 18. A large scale purification was performed as described in Example 24d with the exception that except the soluble lysate was batch absorbed for 3 hours rather than for 1 hour. The final yield was 13 mg pHisBot protein/liter culture. The pHisBot protein represented 0.75% of the total soluble protein.

The above results demonstrate growth conditions under which soluble pHisBot protein is produced (i.e., use of the BL21(DE3)pLysS host, 2X YT medium, 30°C, 1.0 mM IPTG for 5 hours).

b) Optimization Of Purification Parameters

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For optimization of purification conditions, large scale cultures (3 X 1 liter) were grown at 30°C and induced with 1 mM IPTG for 5 hours as described above. The cultures were pooled, distributed to centrifuge bottles, cooled and pelleted as described in Example 24d. The cell pellets were frozen at -70°C until used. Each cell pellet represented 1/3 of a liter starting culture and individual bottles were utilized for each optimization experiment described below. This standardized the input bacteria used for each experiment, such that the yields of affinity purified pHisBot protein could be compared between different optimization experiments.

i) Binding Specificity (pH Protonation)

A lysate of pHisBot culture was prepared in PBS (pH 8.0) and applied to a 3 ml Ni-NTA column equilibrated in PBS (pH 8.0) using a flow rate of 0.2 ml/min (3-4 column volumes/hr) using an Econo chromatography system (BioRad). The column was washed with PBS (pH 8.0) until the absorbance (OD₂₈₀) of the elute was at baseline levels. The flow rate was then increased to 2 ml/min and the column was equilibrated in PBS (pH 7.0). A pH gradient (pH 7.0 to 4.0 in PBS) was applied in order to elute the bound pHisBot protein from the column. Fractions were collected and aliquots were resolved on SDS-PAGE gels. The PAGE gels were subjected to Western blotting and the pHisBot protein was detected using a chicken anti-C. botulinum Type A toxoid antibody as described in Example 22.

From the Western blot analysis it was determined that the pHisBot protein begins to elute from the Ni-NTA column at pH 6.0. This is consistent with the predicted elution of a His-tagged protein monomer at pH 5.9.

These results demonstrate that the pH at which the pHisBot protein is protonated (released) from Ni-NTA resin in PBS buffer is pH 6.0.

ii) Binding Specificity (Imidazole Competition)

In order to define purification conditions under which the native *E. coli* proteins could be removed from the Ni-NTA column while leaving the pHisBot protein bound to the column, the following experiment was performed. A lysate of pHisBot culture was prepared in 50 mM NaHPO₄, 0.5 M NaCl, 8 mM imidazole (pH 7.0). This lysate was applied to a 3 ml Ni-NTA column equilibrated in 50 mM NaHPO₄, 0.5 M NaCl (pH 7.0) using an Econo chromatography system (BioRad). A flow rate of 0.2 ml/min (3-4 column volumes/hr) was utilized. The column was washed with 50 mM NaHPO₄, 0.5 M NaCl (pH 7.0) until the absorbance of the elute returned to baseline. The flow rate was then increased to 2 ml/min.

The column was eluted using an imidazole step gradient [in 50 mM NaHPO₄, 0.5 M NaCl (pH 7.0)]. Elution steps were 20 mM, 40 mM, 60 mM, 80 mM, 100 mM,

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200 mM, 1.0 M imidazole, followed by a wash using 0.1 mM EDTA (to strip the nickel from the column and remove any remaining protein). In each step, the wash was continued until the OD₂₈₀ returned to baseline. Fractions were resolved on SDS-PAGE gels, Western blotted, and pHisBot protein detected using a chicken anti-C. botulinum Type A toxoid antibody as described in Example 22. Duplicate gels were stained with Coomassie blue to detect eluted protein in each fraction.

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The results of the PAGE analysis showed that most of the non-specifically binding bacterial protein was removed by the 20 mM imidiazole wash, with the remaining bacterial proteins being removed in the 40 and 60 mM imidazole washes. The pHisBot protein began to elute at 100 mM imidazole and was quantitatively eluted in 200 mM imidazole.

These results precisely defined the window of imidazole wash stringency that optimally removes *E. coli* proteins from the column while specifically retaining the pHisBot protein in this buffer. These results provided conditions under which the pHisBot protein can be purified free of contaminating host proteins.

iii) Purification Buffers And Optimized Purification Protocols

A variety of purification parameters were tested during the development of an optimized protocol for batch purification of soluble pHisBot protein. The results of these analyses are summarized below.

Batch purifications were performed (as described in Example 24d) using several buffers to determine if alternative buffers could be utilized for binding of the pHisBot protein to the Ni-NTA column. It was determined that quantitative binding of pHisBot protein to the Ni-NTA resin was achieved in either Tris-HCl (pH 7.9) or NaHPO₄ (pH 8.0) buffers. Binding of the pHisBot protein in NaHPO₄ buffer was not inhibited using 5 mM, 8 mM or 60 mM imidazole. Quantitative elution of bound pHisBot protein was obtained in buffers containing 50 mM NaHPO₄, 0.3 M NaCl (pH 3.5-4.0), with or without 10% glycerol. However, quantitation of soluble affinity purified pHisBot protein before and after a freeze thaw (following several weeks storage of the affinity

purified elute at -20°C) revealed that 94% of the protein was recovered using the glycerol-containing buffer, but only 68% of the protein was recovered when the buffer lacking glycerol was employed. This demonstrates that glycerol enhanced the solubility of the pHisBot protein in this low pH buffer when the eluted protein was stored at freezing temperatures (e.g., -20°C). Neutralization of pH by addition of NaH₂PO₄ buffer did not result in obvious protein precipitation.

It was determined that quantitative binding of pHisBot protein using the batch format occurred after 3 hrs (Figure 30), but not after 1 hr of binding at 4°C (the resin was stirred during binding). Figure 30 depicts a Coomaisse blue stained SDS-PAGE gel (7.5% acrylamide) containing samples of proteins isolated during the purification of pHisBot protein from lysate prepared from the BL21(DE3)pLysS host. Each lane was loaded with 5 µl of protein sample mixed with 5 µl of 2X sample buffer and processed as described in Example 22b. Lane 1 contains high molecular weight protein markers (BioRad). Lanes 2 and 3 contain protein eluted from the Ni-NTA resin. Lane 4 contains soluble protein after a 3 hr batch incubation with the Ni-NTA resin. Lanes 5 and 6 contain soluble and total protein, respectively. Figure 30 demonstrates that the pHisBot protein is completely soluble [compare Lanes 5 and 6 which show that a similar amount of the 50 kD pHisBot protein is seen in both; if a substantial amount (greater than 20%) of the pHisBot protein were partially insoluble in the host cell, more pHisBot protein would be seen in lane 6 (total protein) as compared to lane 5 (soluble protein)]. Figure 30 also demonstrates that the pHisBot protein is completely removed from the lysate after batch absorption with the Ni-NTA resin for 3 hours (compare Lanes 4 and 5).

The reported high affinity interaction of the Ni-NTA resin with His-tagged proteins (K_d = 1 x 10⁻¹³ at pH 8.0) suggested that it should be possible to manipulate the resin-protein complexes without significant release of the bound protein. Indeed, it was determined that after the recombinant protein was bound to the Ni-NTA resin, the resin-pHisBot protein complex was highly stable and remained bound following repeated rounds of centrifugation of the resin for 2 min at 1600 x g. When this centrifugation step was performed in a 50 ml tube (Falcon), a tight resin pellet formed.

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This allowed the removal of spent soluble lysate by pouring off the supernatant followed by resuspension of the pellet in wash buffer. Further washes can be performed by centrifugation. The ability to perform additional washes permits the development of protocols for batch absorption of large volumes of lysate with removal of the lysate being performed simply by centrifugation following binding of the recombinant protein to the resin.

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A simplified, integrated purification protocol was developed as follows. A soluble lysate was made by resuspending the induced cell pellet in binding buffer [50 mM NaHPO4, 0.5 M NaCl, 60 mM imidazole (pH 8.0)], sonicating 4 x 20 sec and centrifuging for 20 min at 10,000 x g. NP-40 was added to 0.1% and Ni-NTA resin (equilibrated in binding buffer) was added. Eight milliliters of a 1:1 slurry (resin:binding buffer) was used per liter of starting culture. The mixture was stirred for 3 hrs at 4°C. The slurry was poured into a column having a 1 cm internal diameter (BioRad), washed with binding buffer containing 0.1% NP40, then binding buffer until baseline was established (these steps may alternatively be performed by centrifugation of the resin, resuspension in binding buffer containing NP40 followed by centrifugation and resuspension in binding buffer). Imidazole was removed by washing the resin with 50 mM NaHPO4, 0.3M NaCl (pH 7.0). Protein bound to the resin was eluted using the same buffer (50 mM NaHPO4, 0.3M NaCl) having a reduced pH (pH 3.5-4.0).

A pilot purification was performed following this protocol and yielded 18 mg/liter affinity-purified pHisBot. The pHisBot protein was greater than 90% pure as estimated by Coomassie staining of an SDS-PAGE gel. This represents the highest observed yield of soluble affinity-purified pHisBot protein and this protocol eliminates the need for separate imidazole-containing binding and wash buffers. In addition to providing a simplified and efficient protocol for the affinity purification of recombinant pHisBot protein, the above results provide a variety of purification conditions under which pHisBot protein can be isolated.

EXAMPLE 26

The pHisBot Protein Is An Effective Immunogen

In Example 23 it was demonstrated that neutralizing antibodies are generated in mouse serum after nasal immunization with the pMBot protein. However, the pMBot protein was found to copurify with significant amounts of endotoxin which could not be easily removed. The pHisBot protein, in contrast, could be isolated free of significant endotoxin contamination making pHisBot a superior candidate for vaccine production. To further assess the suitability of pHisBot as a vaccine, the immunogenicity of the pHisBot protein was determined and a comparison of the relative immunogenicity of pMBot and pHisBot proteins in mice was performed as follows.

Two groups of eight BALBc mice were immunized with either pMBot protein or pHisBot protein using Gerbu GMDP adjuvant (CC Biotech). pMBot protein (in PBS containing 10 mM maltose) or pHisBot protein (in 50 mMNaHPO₄, 0.3 M NaCl, 10% glycerol, pH 4.0) was mixed with Gerbu adjuvant and used to immunize mice. Each mouse received an IP injection of 100 µl antigen/adjuvant mix (50 µg antigen plus 1 µg adjuvant) on day 0. Mice were boosted as described above with the exception that the route of administration was IM on day 14 and 28. The mice were bled on day 77 and anti-C. botulinum Type A toxoid titers were determined using serum collected from individual mice in each group (as described in Example 23). The results are shown in Table 41:

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TABLE 41

Anti-C. botulinum Type A Toxoid Serum IgG Titers In Individual Mice Immunized With pMBot or pHisBot Protein

		Prei	mmune'		pMBot²			pHisBot*				
Mouse #		Sample Dilution				Sample Dilution			Sample Dilution			
	1:50	1:250	1:1250	1.6250	1:50	1:250	1:1250	1:6250	_1:50	1:250	1:1250	1:620
1					0.678	0.190	0.055	0.007	1,574	0.799	0.320	0.093
2					1.161	0.931	0.254	0.075	1.513	0.829	0.409	0.134
3					1.364	0.458	0.195	0.041	1.596	1.028	0.453	0.122
4					1.622	1.189	0.334	0.067	1.552	0.840	0.348	0.090
5					1.612	1.030	0.289	0.067	1.629	1.580	0.895	0.233
6					0.913	0.242	0.069	0.013	1.485	0.952	0.477	0.145
7					0.910	0.235	0.058	0.014	1.524	0.725	0.269	0.069
8					0.747	0.234	0.058	, 0.014	1.274	0.427	0.116	0.029
Mean Titer	0.048	0.021	0.011	0.002	1.133	0.564	0.164	0.037	1.518	0.896	0.411	0.114

The preimmune sample represents the average from 2 sets of duplicate wells containing serum from a individual mouse immunized with recombinant *Staphylococcus* enterotoxin B (SEB) antigen. This antigen is immunologically unrelated to *C. botulinum* toxin and provides a control serum.

Average of duplicate wells.

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The results shown above in Table 41 demonstrate that both the pMBot and pHisBot proteins are immunogenic in mice as 100% of the mice (8/8) in each group seroconverted from non-immune to immune status. The results also show that the average titer of anti-C. botulinum Type A toxoid IgG is 2-3 fold higher after immunization with the pHisBot protein relative to immunization with the pMBot protein. This suggests that the pHisBot protein may be a superior immunogen to the pMBot protein.

EXAMPLE 27

Immunization With The Recombinant pHisBot Protein Generates Neutralizing Antibodies

The results shown in Example 26 demonstrated that both the pHisBot and pMBot proteins were capable of inducing high titers of anti-C. botulinum type A toxoid-reactive antibodies in immunized hosts. The ability of the immune sera from mice immunized with either the pHisBot or pMBot proteins to neutralize C. botulinum type A toxoid in vivo was determined using the mouse neutralization assay described in Example 23b.

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The two groups of eight BALBc mice immunized with either pMBot protein or pHisBot protein in Example 26 were boosted again one week after the bleeding on day 77. The boost was performed by mixing pMBot protein (in PBS containing 10 mM maltose) or pHisBot protein (in 50 mM NaHPO₄, 0.3 M NaCl, 10% glycerol, pH 4.0) with Gerbu adjuvant as described in Example 26. Each mouse received an IP injection of 100 µl antigen/adjuvant mix (50 µg antigen plus 1 µg adjuvant). The mice were bled 6 days after this boost and the serum from mice within a group was pooled. Serum from preimmune mice was also collected (this serum is the same serum described in the footnote to Table 41).

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The presence of neutralizing antibodies in the pooled or preimmune serum was detected by challenging mice with 5 LD₅₀ units of type A toxin mixed with 100 µl of pooled serum. The challenge was performed by mixing (per mouse to be injected) 100 µl of serum from each pool with 100 µl of purified type A toxin standard (50 LD₅₀ /ml prepared as described in Example 23b) and 500 µl of gel-phosphate. The mixtures were incubated for 30 min at room temperature with occasional mixing. Each of four mice were injected IP with the mixtures (0.7 ml/mouse). The mice were observed for signs of botulism for 72 hours. Mice receiving toxin mixed with serum from mice immunized with either the pHisBot or pMBot proteins showed no signs of botulism intoxication. In contrast, mice receiving preimmune serum died in less than 24 hours.

These results demonstrate that antibodies capable of neutralizing C. botulinum type A toxin are induced when either of the recombinant C. botulinum C fragment proteins pHisBot or pMBot are used as immunogens.

EXAMPLE 28

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Cloning And Expression Of The C Fragment of C. botulinum Serotype A Toxin In E. coli Utilizing A Native Gene Fragment

In Example 22 above, a synthetic gene was used to express the C fragment of C. botulinum serotype A toxin in E. coli. The synthetic gene replaced non-preferred (i.e., rare) codons present in the C fragment gene with codons which are preferred by E. coli. The synthetic gene was generated because it was been reported that genes which have a high A/T content (such as most clostridial genes) creates expression difficulties in E. coli and yeast. Furthermore, LaPenotiere et al. suggested that problems encountered with the stability (non-fusion constructs) and solubility (MBP fusion constructs) of the C fragment of C. botulinum serotype A toxin when expressed in E. coli was most likely due to the extreme A/T richness of the native C. botulinum serotype A toxin gene sequences (LaPenotiere, et al., supra).

In this example, it was demonstrated that successful expression of the C fragment of C. botulinum type A toxin gene in E. coli does not require the elimination of rare codons (i.e., there is no need to use a synthetic gene). This example involved a) the cloning of the native C fragment of the C. botulinum serotype A toxin gene and construction of an expression vector and b) a comparison of the expression and purification yields of C. botulinum serotype A C fragments derived from native and synthetic expression vectors.

a) Cloning Of The Native C Fragment Of The C. botulinum Serotype A Toxin Gene And Construction Of An Expression Vector

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The serotype A toxin gene was cloned from *C. botulinum* genomic DNA using PCR amplification. The following primer pair was employed: 5'-CGCCATGGCTAG ATTATTATCTACATTTAC-3' (5' primer, *Nco*I site underlined; SEQ ID NO:29) and 5'-GCAAGCTTCTTGACAGACTCATGTAG-3' (3' primer, *HindIII* site underlined; SEQ ID NO:30). *C. botulinum* type A strain was obtained from the American Type Culture Collection (ATCC#19397) and grown under anaerobic conditions in Terrific broth medium. High molecular-weight *C. botulinum* DNA was isolated as described in Example 11. The integrity and yield of genomic DNA was assessed by comparison with a serial dilution of uncut lambda DNA after electrophoresis on an agarose gel.

The gene fragment was cloned by PCR utilizing a proofreading thermostable DNA polymerase (native Pfu polymerase). PCR amplification was performed using the above primer pair in a 50µl reaction containing 10mM Tris-HCl (pH 8.3), 50mM KCl, 1.5mM MgCl₂, 200µM each dNTP, 0.2µM each primer, and 50ng C. botulinum genomic DNA. Reactions were overlaid with 100µl mineral oil, heated to 94°C 4 min, 0.5µl native Pfu polymerase (Stratagene) was added, and thirty cycles comprising 94°C for 1 min, 50°C for 2 min, 72°C for 2 min were carried out followed by 10 min at 72°C. An aliquot (10µl) of the reaction mixture was resolved on an agarose gel and the amplified native C fragment gene was gel purified using the Prep-A-Gene kit (BioRad) and ligated to pCRScript vector DNA (Stratagene). Recombinant clones were isolated and confirmed by restriction digestion, using standard recombinant molecular biology techniques [Sambrook et al. (1989), supra]. In addition, the sequence of approximately 300 bases located at the 5' end of the C fragment coding region were obtained using standard DNA sequencing methods. The sequence obtained was identical to that of the published sequence.

An expression vector containing the native C. botulinum serotype A C fragment gene was created by ligation of the Ncol-HindIII fragment containing the C fragment

gene from the pCRScript clone to *NheI-HindIII* restricted pETHisa vector (Example 18b). The *NcoI* and *NheI* sites were filled in using the Klenow enzyme prior to ligation; these sites were thus blunt-end ligated together. The resulting construct was termed pHisBotA (native). pHisBotA (native) expresses the *C. botulinum* serotype A C fragment with a his-tagged N terminal extension which has the following sequence: MetGlyHisHisHisHisHisHisHisHisHisHisHisSerSerGlyHis*IleGluGlyArg*HisMetAla (SEQ ID NO:24), where the underlining represents amino acids encoded by the *C. botulinum* C fragment gene (this N terminal extension contains the recognition site for FactorXa protease, shown in italics, which can be employed to removed the polyhistdine tract from the N-terminus of the fusion protein). The pHisBot (native) construct expresses the identical protein as the pHisBot construct (Ex. 24c; herein after the pHisBotA) which contains the synthetic gene.

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The predicted DNA sequence encoding the native *C. botulinum* serotype A C fragment gene contained within pHisBotA (native) is listed in SEQ ID NO:31 [the start of translation (ATG) is located at nucleotides 108-110 and the stop of translation (TAA) is located at nucleotides1494-1496 in SEQ ID NO:31] and the corresponding amino acid sequence is listed in SEQ ID NO:26 (*i.e.*, the same amino acid sequence as that produced by pHisBotA containing synthetic gene sequences).

b) Comparison Of The Expression And Purification
Yields Of C. botulinum Serotype A C Fragments
Derived From Native And Synthetic Expression
Vectors

Recombinant plasmids containing either the native or the synthetic C. botulinum serotype A C fragment genes were transformed into E. coli strain Bl21(DE3) pLysS and protein expression was induced in 1 liter shaker flask cultures. Total protein extracts were isolated, resolved on SDS-PAGE gels and C. botulinum C fragment protein was identified by Western analysis utilizing a chicken anti-C. botulinum serotype A toxoid antiserum as described in Example 22.

Briefly, 1 liter (2XYT + 100 μ g/ml ampicillin and 34 μ g/ml chloramphenicol) cultures of bacteria harboring either the pHisBotA (synthetic) or pHisBotA (native) plasmids in the Bl21(DE3) pLysS strain were induced to express recombinant protein by addition of IPTG to 1mM. Cultures were grown at 30-32°C, IPTG was added when the cell density reached an OD₆₀₀ 0.5-1.0 and the induced protein was allowed to accumulate for 3-4 hrs after induction.

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The cells were cooled for 15 min in a ice water bath and then centrifuged for 10 min at 5000 rpm in a JA10 rotor (Beckman) at 4°C. The cell pellets were resuspended in a total volume of 40 mls 1X binding buffer (40 mM imidazole, 0.5 M NaCl, 50 mM NaPO₄, pH 8.0), transferred to two 50 ml Oakridge tubes and frozen at -70°C for at least 1 hr. The tubes were then thawed and the cells were lysed by sonication (using four successive 20 second bursts) on ice. The suspension was clarified by centrifugation 20-30 min at 9,000 rpm (10,000g) in a JA-17 rotor. The soluble lysate was batch absorbed to 7 ml of a 1:1 slurry of NiNTA resin:binding buffer by stirring 2-4 hr at 4°C. The slurry was centrifuged for 1 min at 500g in 50 ml tube (Falcon), resuspended in 5 mls binding buffer and poured into a 2.5 cm diameter column (BioRad). The column was attached to a UV monitor (ISCO) and the column was washed with binding buffer until a baseline was established. Imidazole was removed by washing with 50mM NaPO₄, 0.3 M NaCl, 10% glycerol, pH 7.0 and bound protein was eluted using 50mM NaPO₄, 0.3 M NaCl, 10% glycerol, pH 3.5-4.0.

The eluted proteins were stored at 4°C. Samples of total, soluble, and eluted proteins were resolved by SDS-PAGE. Protein samples were prepared for electrophoresis by mixing 1µl total (T) or soluble (S) protein with 4 µl PBS and 5 µl 2X SDS-PAGE sample buffer, or 5 µl eluted (E) protein and 5 µl 2X SDS-PAGE sample buffer. The samples were heated to 95°C for 5 min, then cooled and 5 or 10 µls were loaded on 12.5% SDS-PAGE gels. Broad range molecular weight protein markers (BioRad) were also loaded to allow the MW of the identified fusion proteins to be estimated. After electrophoresis, protein was detected either generally by

staining gels with Coomassie blue, or specifically, by blotting to nitrocellulose for Western blot detection of specific immunoreactive protein.

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For Western blot analysis, the gels were blotted, and protein transfer was confirmed by Ponceau S staining as described in Example 22. After blocking the blots for 1 hr at room temperature in blocking buffer (PBST and 5% milk), 10 ml of a 1/500 dilution of an anti-C. botulinum toxin A IgY PEG prep (Ex. 3) in blocking buffer was added and the blots were incubated for an additional hour at room temperature. The blots were washed and developed using a rabbit anti-chicken alkaline phosphatase conjugate (Boehringer Mannheim) as the secondary antibody as described in Ex. 22. This analysis detected C. botulinum toxin A-reactive proteins in the pHisBotA (native and synthetic) protein samples (corresponding to the predicted full length proteins identified by Coomassie staining).

A gel containing proteins expressed from the pHisBot and pHisBot (native) constructs during various stages of purification and stained with Coomassie blue is shown in Figure 31. In Figure 31, lanes 1-4 and 9 contain proteins expressed by the pHisBotA construct (i.e., the synthetic gene) and lanes 5-8 contain proteins expressed by the pHisBotA (native) construct. Lanes 1 and 5 contain total protein extracts; lanes 2 and 6 contain soluble protein extracts; lanes 3 and 7 contain proteins which flowed through the NiNTA columns; lanes 4, 8 and 9 contain protein eluted from the NiNTA columns and lane 10 contains molecular weight markers.

The above purification resulted in a yield of 3 mg (native gene) or 11 mg (synthetic gene) of affinity purified protein from a 1 liter starting culture, of which at least 90-95% of the protein was a single band of the predicted MW (50kd) and immunoreactivity for recombinant *C. botulinum* serotype A C fragment protein. Other than the level of expression, no difference was observed between the native and the synthetic gene expression systems.

These results demonstrate that soluble C. botulinum serotype A C fragment protein can be expressed in E. coli and purified utilizing either native or synthetic gene sequences.

EXAMPLE 29

Generation Of Neutralizing Antibodies Using A Recombinant

C. botulinum Serotype A C Fragment Protein Containing A Six Residue His-Tag

In Example 27, neutralizing antibodies were generated utilizing the pHisBotA protein, which contains a histidine-tagged N-terminal extension comprising 10 histidine residues. To determine if the generation of neutralizing antibodies is dependent on the presence of this particular his-tag, a protein containing a shorter N-terminal extension (comprising 6 histidine residues) was produced and tested for the ability to generate neutralizing antibodies. This example involved a) the cloning and expression of the p6HisBotA(syn) protein and b) the generation and characterization of hyperimmune serum.

a) Cloning And Expression Of The p6HisBotA(syn) Protein

The p6HisBotA(syn) construct was generated as described below; the term "syn" designates the presence of synthetic gene sequences. This construct expresses the C frgament of the C. botulinum serotype A toxin with a histidine-tagged N terminal extension having the following sequence: MetHisHisHisHisHisHisHisMetAla (SEQ ID NO:32); the amino acids encoded by the botulinal C fragment gene are underlined and the vector encoded amino acids are presented in plain type.

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6XHis oligonucleotides [5'-TATGCATCACCATCACCATCA-3' (SEQ ID NO:33) and 5'-CATGTGATGGTGATGGTGATGGTGATGCA-3' (SEQ ID NO:34) were annealed as follows. One microgram of each oligonucleotide was mixed in total of 20 μl 1X reaction buffer 2 (NEB) and the mixture was heated at 70°C for 5 min and then incubated at 42°C for 5 min. The annealed oligonucleotides were then ligated with gel purified Ndel/HindIII cleaved pET23b (T7 promoter) or pET21b (T7lac promoter) DNA and the gel purified Ncol/HindIII C. botulinum serotype A C fragment synthetic gene fragment derived from pAlterBot (Ex. 22). Recombinant clones were isolated and confirmed by restriction digestion. The DNA sequence encoding the 6X his-

tagged BotA protein contained within p6HisBotA(syn) is listed in SEQ ID NO:35. The amino acid sequence of the p6XHisBotA protein is listed in SEQ ID NO:36.

The resulting recombinant p6XHisBotA plasmid was transformed into the BL21(DE3) pLysS strain, and 1 liter cultures were grown, induced and harvested as described in Example 28. His-tagged protein was purified as described in Example 28, with the following modifications. The binding buffer (BB) contained 5 mM imidazole rather than 40 mM imidazole and NP40 was added to the soluble lysate to a final concentration of 0.1%. The bound material was washed on the column with BB until the baseline was established, then the column was washed successively with BB+20 mM imidazole and BB+40 mM imidazole. The column was eluted as described in Example 28.

In the case of the pET23-derived expression system, high level expression of insoluble 6HisBotA protein was induced. The pET21-derived vector expressed lower levels of soluble protein that bound the NiNTA resin and eluted in the 40 mM imidazole wash rather than during the low pH elution. These results (i.e., low level expression of a soluble protein) are consistent with the results obtained with pHisBotA protein (Ex. 25); the pHisBotA construct, like the pET21-derived vector, contains the T7lac rather than T7 promoter.

The 6HisBotA protein thus elutes under less stringent conditions than the 10X histidine-containing pHisBot protein (100-200 mM imidazole; Ex. 25) presumably due to the reduction in the length of the his-tag. The eluted protein was of the predicted size [i.e., slightly reduced in comparison to pHisBotA protein].

b) Generation And Characterization Of Hyperimmune Serum

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Eight BALBc mice were immunized with purified 6HisBotA protein using Gerbu GMDP adjuvant (CC Biotech). The 40 mM imidazole elution was mixed with Gerbu adjuvant and used to immunize mice. Each mouse received a subcutaneous injection of 100 μl antigen/adjuvant mix (12 μg antigen + 1 μg adjuvant) on day 0. Mice were subcutaneously boosted as above on day 14 and bled on day 28. Control

mice received pHisBotB protein (prepared as described in Ex. 35 below) in Gerbu adjuvant.

Anti-C. botulinum serotype A toxoid titers were determined in serum from individual mice from each group using the ELISA described in Example 23a with the exception that the initial testing serum dilution was 1:100 in blocking buffer containing 0.5% Tween 20, followed by serial 5-fold dilutions into this buffer. The results of the ELISA demonstrated that seroconversion (relative to control mice) occurred in all 8 mice.

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The ability of the anti-C. botulinum serotype A C fragment antibodies present in serum from the immunized mice to neutralize native C. botulinum type A toxin was tested using the mouse neutralization assay described in Example 23b. The amount of neutralizing antibodies present in the serum of the immunized mice was determined using serum antibody titrations. The various serum dilutions (0.01 ml) were mixed with 5 LD₅₀ units of C. botulinum type A toxin and the mixtures were injected IP into mice. The neutralizations were performed in duplicate. The mice were then observed for signs of botulism for 4 days. Undiluted serum was found to protect 100% of the injected mice while the 1:10 diluted serum did not. This corresponds to a neutralization titer of 0.05-0.5 IU/ml.

These results demonstrate that neutralizing antibodies were induced when the 6HisBotA protein was utilized as the immunogen. Furthermore, these results demonstrate that seroconversion and the generation of neutralizing antibodies does not depend on the specific N terminal extension present on the recombinant C. botulinum type A C fragment proteins.

EXAMPLE 30

Construction Of Vectors For The Expression Of His-Tagged

C. botulinum Type A Toxin C Fragment Protein Using the Synthetic Gene

A number of expression vectors were constructed which contained the synthetic C. botulinum type A toxin C fragment gene. These constructs vary as to the promoter (T7 or T7lac) and repressor elements (lacIq) present on the plasmid. The T7 promoter is a stronger promoter than is the T7lac promoter. The various constructs provide varying expression levels and varying levels of plasmid stability. This example involved a) the construction of expression vectors containing the synthetic C. botulinum type A C fragment gene and b) the determination of the expression level achieved using plasmids containing either the kanamycin resistance or the ampicillin resistance genes in small scale cultures.

a) Construction Of Expression Vectors Containing The Synthetic C. botulinum Type A C Fragment Gene

Expression vectors containing the synthetic *C. botulinum* type A C fragment gene were engineered to utilize the kanamycin resistance rather than the ampicillin resistance gene. This was done for several reasons including concerns regarding the presence of residual ampicillin in recombinant protein derived from plasmids containing the ampicillin resistance gene. In addition, ampicillin resistant plasmids are more difficult to maintain in culture; the β-lactamase secreted by cells containing ampicillin resistant plasmids rapidly degrades extracellular ampicillin, allowing the growth of plasmid-negative cells.

A second altered feature of the expression vectors is the inclusion of lacIq gene in the plasmid. This repressor lowers expression from lac regulated promoters (the chromosomally located, lactose regulated T7 polymerase gene and the plasmid located T7lac promoter). This down regulates uninduced protein expression and can enhance the stability of recombinant cell lines. The final alteration to the vectors is the

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inclusion of either the T7 or T7lac promoters that drive high or moderate level expression of recombinant protein, respectively.

The expression plasmids were constructed as follows. In all cases, the protein expressed is the pHisBotA(syn) protein previously described, and the only differences between constructs is the alteration of the various regulatory elements described above.

i) Construction Of pHisBotA(syn) kanT7lac

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The pHisBotA(syn) kan T7lac construct was made by inserting the Sapl/Xhol fragment containing the C. botulinum type A C fragment from pHisBotA(syn) into pET24 digested with Sapl/Xhol (Novagen; fragment contains kan gene and origin of replication). The desired construct was selected for kanamycin resistance and confirmed by restriction digestion.

ii) Construction Of pHisBotA(syn) kan lacIq T7lac

The pHisBotA(syn) kan lacIq T7lac construct was made by inserting the Xbal/HindIII fragment containing the C. botulinum type A C fragment from pHisBotA(syn)kanT7lac into the pET24a vector digested with Xbal/HindIII. The resulting construct was confirmed by restriction digestion.

iii) Construction Of pHisBotA(syn) kan lacIq T7

The pHisBotA(syn) kan lacIq T7 construct was made by inserting the XbaI/HindIII fragment containing the C. botulinum type A C fragment from pHisBotA(syn) kan lacIq T7lac into XbaI/HindIII-digested pHisBotB(syn) kan lacIq T7 (described in Ex 37c below). The resulting construct was confirmed by restriction digestion.

b) Determination Of The Expression Level Achieved
Using Plasmids Containing Either The Kanamycin
Resistance Or The Ampicillin Resistance Genes In
Small Scale Cultures

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One liter cultures of pHisBotA(syn) kan T7lac/Bl21(DE3)pLysS and pHisBotA(syn) amp T7lac/Bl21(DE3)pLysS [this is the previously designated pHisBotA(syn) construct] were grown, induced and his-tagged proteins were purified as described in Example 28. No differences in yield or protein integrity/purity were observed.

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These results demonstrate that the antigen induction levels from expression constructs were not affected by the choice of ampicillin versus kanamycin antibiotic resistance genes.

EXAMPLE 31

Fermentation Of Cells Expressing Recombinant Botulinal Proteins

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a) Fermentation Culture Of Cells Expressing Recombinant Botulinal Proteins

Fermentation cultures were grown under the following conditions which were optimized for growth of the BL21(DE3) strains containing pET derived expression vectors. An overnight 1 liter feeder culture was prepared by inoculating of 1 liter media (in a 2L shaker flask) with a fresh colony grown on an LB kan plate. The feeder culture contained: 600 mls nitrogen source [20 gm yeast extract (BBL) and 40 gm tryptone (BBL)/600 mls], 200 mls 5X fermentation salts (per liter: 48.5 gm K₂HPO₄, 12 gm NaH₂PO₄•H₂O, 5 gm NH₄Cl, 2.5 gm NaCl), 180 mls dH₂O, 20 mls 20% glucose, 2 mls 1 M MgSO₄, 5 mls 0.05M CaCl₂ and 4 mls of a 10 mg/ml kanamycin stock. All solutions were sterilized by autoclaving, except the kanamycin stock which was filter sterilized.

An aliquot (5 ml) of the feeder culture broth was removed prior to inoculation, and grown for 2 days at 37°C as a culture broth sterility control. Growth was not observed in this control culture in any of the fermentations performed.

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The inoculated feeder culture was grown for 12-15 hrs (ON) at 30-37°C. Care was taken to prevent oversaturation of this culture. The saturated feeder culture was added to 10L of fermentation media in fermenter (BiofloIV, New Brunswick Scientific, Edison, NJ) as follows. The fermenter was sterilized 120 min at 121°C with dH₂O. The sterile water was removed, and fermentation media added as follows: 6 liters nitrogen source, 2 liters 5X fermentation salts, 2 liters 2% glucose, 20 mls 1 M MgSO₄, 50 mls 0.05 M CaCl₂, 2.5-3.5 mls Macol P 400 antifoam (PPG Industries Inc., Gurnee, IL), 40 mls 10mg/ml kanamycin and 10 mls trace elements (8 gm FeSO₄•7H₂O, 2 gm MnSO₄•H₂O, 2 gm AlCl₃•6H₂O, 0.8 gm CoCl•6H₂O, 0.4 gm ZnSO₄•7H₂O, 0.4 gm Na₂MoO₄•2H₂O, 0.2 gm CuCl₂•2H₂O, 0.2 gm NiCl₂, 0.1 gm H₃BO₄/200mls 5 M HCl). All solutions were sterilized by autoclaving, except the kanamycin stock which was filter sterilized. Fermentation media was prewarmed to 37°C before the addition of the feeder culture.

After the addition of the feeder culture, the culture was fermented at 37°C, 400 rpm agitation, and 10 l/min air sparging. The DO₂ control was set to 20% PID and dissolved oxygen levels were controlled by increasing the rate of agitation from 400-850 rpm under DO₂ control. DO₂ levels were maintained at greater than or equal to 20% throughout the entire fermentation. When agitation levels reached 500-600 rpm the temperature was lowered to 30°C to reduce the oxygen consumption rate. Culture growth was continued until endogenous carbon sources were depleted. In these fermentations, glucose was depleted first [monitored with a glucose monitoring kit (Sigma)], followed by assimilation of acetate and other acidic carbons [monitored using an acetate test kit (Boehringer Mannheim)]. During the assimilation phase, the pH rose from 6.6-6.8 (starting pH) to 7.4-7.5, at which time the bulk of the remaining carbon source was depleted. This was signaled by a drop in agitation rate (from a maximum of 700-800 rpm) and a rise in DO₂ levels >30%. This corresponds to a OD₆₀₀ reading of 18-20/ml. At this point a fed batch mode was initiated, in which a

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feed solution of 50% glucose was added at a rate of approximately 4 gm glucose/liter/hr. The pH was adjusted to 7.0 by the addition of 25% H₃PO₄ (approximately 60 mls). Culture growth was continued and reached peak oxygen consumption within the next 3 hrs of growth (while the remaining residual non-glucose carbon sources were assimilated). This phase is characterized by a slow increase in pH, and air sparging was increased to 15L/min, to keep the maximum rpm below 850.

Once the residual acidic carbon sources are depleted the agitation rate decreases to 650-750 rpm and the pH begins to drop. pH control was maintained at 7.0 PID by regulated pump addition of a sterile 4M NaOH solution which was consumed at a steady rate for the remainder of the fermentation. Growth was continued at 30°C, and the cultures were grown linearly at a growth rate of 4-7 OD₆₀₀ units/hr, to at least 81.5 OD₆₀₀ units/ml (>30g/l dry cell weight) without induction. Antifoam (a 1:1 dilution with filter sterilized 100% ethanol) was added as necessary throughout the fermentation to prevent foaming.

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During the fed batch mode, glucose was assimilated immediately (concentration in media consistently less than 0.1 gm/liter) and acetate was not produced in significant levels by the pET plasmid/BL21(DE3) cell lines tested (approximately 1 gm/liter at end of fermentation; this is lower than that observed in harvests from shaker flask cultures utilizing the same strains). This was fortuitous, since high levels of acetate has been shown to inhibit induction levels in a variety of expression systems. The above described conditions were found to be highly reproducible between fermentations and utilizing different expression plasmids. As a result, glucose and acetate level monitoring were no longer preformed during fermentation.

b) Induction Of Fermentation Cultures

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Induction with IPTG (250 mg-10 gms, depending on the expression vector and experiment) was initiated 1-3 hrs after initiation of the glucose feed (30-50 OD₆₀₀/ml). The growth rate after induction was monitored on a hourly basis. Aliquots (5-10 ml) of cells were harvested at the time of induction, and at hourly intervals post-induction. Optical density readings were determined by measuring the absorbance at 600 nm of

10 μl culture in 990 μl PBS versus a PBS control. The growth rate after induction was found to vary depending on the expression system utilized.

c) Monitoring Of Fermentation Cultures

Fermentation cultures were monitored using the following control assays.

i) Colony Forming Ability

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An aliquots of cells were removed from the cultures at each timepoint sampled (uninduced and at various times after induction) were serially diluted in PBS (dilution $1=15 \mu l$ cells/3 ml PBS, dilution $2=15 \mu l$ of dilution 1/3 ml PBS, dilution 3=3 or 6 μl of dilution 2/3mls PBS) and $100 \mu l$ of dilution 3 was plated on an LB or TSA (trypticase soy agar) plate. The plates were incubated ON at 37° C and then the colonies are counted and scored for macro or micro growth.

ii) Phenotypic Characterization

Colonies growing on LB or TSA plates (above) from uninduced and induced timepoints were replica plated onto LB+kan, LB+chloramphenicol (for fermentations utilizing LysS or pACYCGro plasmids), LB+kan+1mM IPTG and LB plates, in this order. The plates were grown 6-8 hrs at 37°C and growth was scored on each plate for a minimum of 40-50 well isolated colonies. The percentage of cells retaining the plasmid at time of induction (*i.e.*, uninduced cultures immediately prior to the addition of IPTG) was determined to be the # colonies LB+Kan (or chloramphenicol) plate/# colonies LB plate X 100%. The percentage of cells with mutated pET plasmids was determined to be the # colonies LB+Kan+IPTG plate/# colonies LB plate X 100%. Colonies on all LB plates were scored morphologically for *E. coli* phenotype as a contamination control. Morphologically detectable contaminant colonies were not detected in any fermentation.

Recombinant BotA Protein Induction iii)

A total of 10 OD₆₀₀ units of cells (e.g., 200 μ l of cells at OD₆₀₀=50/ml) were removed from each timepoint sample to a 1.5 ml microfuge tube and pelleted for 2 min at maximum rpm in a microfuge. The pellets were resuspended in 1 ml of 50 mM NaHPO₄, 0.5 M NaCl, 40mM imidazole buffer (pH 6.8) containing 1 mg/ml lysozyme. The samples were incubated for 20 min at room temperature and stored ON at -70°C. Samples were thawed completely at room temperature and sonicated 2 X 10 seconds with a Branson Sonifier 450 microtip probe at # 3 power setting. The samples were centrifuged for 5 min at maximum rpm in a microfuge.

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An aliquot (20 µl) of the protein samples were removed to 20 µl 2X sample buffer, before or after centrifugation, for total and soluble protein extracts, respectively. The samples were heated to 95°C for 5 min, then cooled and 5 or 10 µl were loaded onto 12.5% SDS-PAGE gels. High molecular weight protein markers (BioRad) were also loaded to allow for estimation of the MW of identified fusion proteins. After electrophoresis, protein was detected either generally by staining gels with Coomassie blue, or specifically, by blotting onto nitrocellulose (as described in Ex. 28) for Western blot detection of specific his-tagged proteins utilizing a NiNTA-alkaline phosphatase conjugate exactly as described by the manufacturer (Qiagen).

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Recombinant Antigen Purification iv)

At the end of each fermentation run, 1-10 liters of culture were harvested from the fermenter and the bacterial cells were pelleted by centrifugation at 6000 rpm for 10 min in a JA10 rotor (Beckman). The cell pellets were stored frozen at -70°C or utilized immediately without freezing. Cell pellets were resuspended to 15-20% weight to volume in resuspension buffer (generally 50 mM NaPO₄, 0.5 M NaCl, 40mM imidazole, pH 6.8) and lysed utilizing either sonication or high pressure homogenization.

For sonication, the resuspension buffer was supplemented with lysozyme to 1 mg/ml, and the suspension was incubated for 20 min. at room temp. The sample was then frozen ON at -70°C, thawed and sonicated 4 X 20 seconds at microtip maximum to reduce viscosity. For homogenization, the cells were lyzed by 2 passes through a homogenizer (Rannie Mini-lab type 8.30 H) at 600 Bar. Cell lysates were clarified by centrifugation for 30 min at 10,000 rpm in a JA10 rotor.

For IDA chromatography, samples were flocculated utilizing polyethyleneimine (PEI) prior to centrifugation. Cell pellets were resuspended in cell resuspension buffer (CRB: 50 mM NaPO₄, 0.5 M NaCl, 40 mM imidazole, pH 6.8) to create a 20% cell suspension (wet weight of cells/volume of CRB) and cell lysates were prepared as described above (sonication or homogenization). PEI (a 2% solution in dH₂O, pH 7.5 with HCl) was added to the cell lysate a final concentration of 0.2%, and stirred for 20 min at room temperature prior to centrifugation (8,500 rpm in JA10 rotor for 30 minutes at 4°C). This treatment removed RNA, DNA and cell wall components, resulting in a clarified, low viscosity lysate ("PEI clarified lysate").

His-tagged proteins were purified from soluble lysates by metal-chelate affinity chromatography using either a NiNTA resin (as described in Ex. 28) or an IDA (iminodiacetic acid) resin as described below.

IDA resin affinity purifications were performed utilizing a low pressure chromatography system (ISCO). A 7 ml (small scale) or 70 ml (large scale) Chelating Sepharose Fast Flow (Pharmacia) affinity column was poured; in addition, a second guard column was poured and attached in line with the first column (to capture Ni ions that leached off the affinity column). The columns were washed with 3 column volumes of dH₂O. The guard column was then removed and the affinity column was washed with 0.3 M NiSO₄ until resistivity was established, then with dH₂O until the resistivity returned to baseline. The columns were reconnected and equilibrated with cell resuspension buffer (CRB; 50 mM NaPO₄, 0.5 M NaCl, 40 mM imidazole, pH 6.8). The clarified sample (in CRB) was loaded. Flow rates were 5 ml/min for small scale columns and 20 ml/min for large scale columns. After sample loading, the column was washed with CRB until a baseline established and bound protein was eluted with elution buffer (50 mM NaPO₄, 0.5 M NaCl, 800 mM imidazole, 20% glycerol, pH 6.8 or 8.0). Protein samples were stored at 4°C or -20°C. The yield of

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eluted protein was established by measuring the OD₂₈₀ of the elutions, with a 1 mg/ml solution of protein assumed to yield an absorbance reading of 2.0.

The IDA columns may be regenerated and reused multiple times (>10). To regenerate the column, the column was washed with 2-3 column volumes of H₂O, then 0.05 M EDTA until all of the blue/green color was removed followed by a wash with dH₂O. The IDA columns were sterilized with 0.1 M NaOH (using at least 3 column volumes but not more than 50 minutes contact time with column packing material), then washed with 3 column volumes 0.05 M NaPO₄, pH 5.0, then dH₂O and stored at room temperature in 20 % ethanol.

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EXAMPLE 32

Construction Of A Folding Chaperone Overexpression System

Co-overexpression of the *E. coli* GroEL/GroES folding chaperones in a cell expressing a recombinant foreign protein has been reported to enhance the solubility of some foreign proteins that are otherwise insoluble when expressed in *E. coli* [Gragerouu *et al.* (1992) Proc. Natl. Acad. Sci. USA 89:10344]. The improvement in solubility is thought to be due to chaperone-mediated binding and unfolding of insoluble denatured proteins, thus allowing multiple attempts for productive refolding of recombinant proteins. By overexpressing the chaperones, the unfolding/refolding reaction is driven by excess chaperone, resulting, in some cases, in higher yields of soluble protein.

In this example, a chaperone overexpression system, compatible with pET vector expression systems, was constructed to facilitate testing chaperone-mediated solubilization of *C. botulinum* type A proteins. This example involved the cloning of the GroEL/ES operon and construction of a pLysS-based chaperone hyperexpression system.

The GroEL/GroES operon was PCR amplified and cloned into the pCRScript vector as described in Example 28. The following primer pair was used: 5'-CGCAT ATGAATATTCGTCCATTGCATG-3' (SEQ ID NO:37) [5' primer, start codon of

groES gene converted to NdeI site (underlined)] and 5'-GGAAGCTTGCAGGGCAAT TACATCATG (SEQ ID NO:38) (3' primer, stop codon of groEL gene italicized, engineered HindIII site underlined). Following amplification, the chaperone operon was excised as an NdeI/HindIII fragment and cloned into pET23b digested with NdeI and HindIII. This construction places the Gro operon under the control of the T7 promoter of the pET23 vector. The desired construct was confirmed by restriction digestion.

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The T7 promoter-Gro operon-T7 terminator expression cassette was then excised as a *BgIII/BspEI* (filled) fragment and cloned into *BamHI* (compatible with *BgIII)/HindIII* (filled) cleaved pLysS plasmid (this removed the T7 lysozyme gene). The resulting construct was designated pACYCGro, since the plasmid utilizing the pACYC184 origin from the plysS plasmid. Proper construction was confirmed by restriction digestion.

pACYCGro was transformed into BL21(DE3), cultures were grown and induced with 1 mM IPTG as described in preceding examples. Total and soluble protein extracts were generated from cells removed before and after IPTG induction and were resolved on a 12.5 % SDS-PAGE gel and stained with Coomassie blue. This analysis revealed that high levels of soluble GroEl and GroES proteins were made in the induced cells. These results demonstrated that the chaperone hyper-expression system was functional.

EXAMPLE 33

Growth Of BotA/pACYCGro Cell Lines In Fermentation Cultures

Induction of BL21(DE3) cells lacking the LysS plasmid which contained BotA expression constructs grown in shaker flask or fermentation culture resulted in the expression of primarily insoluble BotA protein. Fermentation cultures were performed to determine if the simultaneous overexpression of the Gro operon and recombinant C. botulinum type A proteins (BotA proteins) resulted in enhanced solubility of the

recombinant BotA protein. This example involved the fermentation of pHisBotA(syn)kan lacIq T7lac/pACYCGro BL21(DE3) and pHisBotA(syn)kan lacIq T7/pACYCGro BL21(DE3) cell lines. The fermentations were repeated exactly as described in Example 31. Chloramphenicol (34 µg/ml) was included in the feeder and fermentation cultures.

a) Fermentation Of pHisBotA(syn)kan lacIq T7lac/pACYCGro BL21(DE3) Cells

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For fermentation of cells containing plasmids comprising the T7lac promoter, induction was with 2 gms IPTG at 1 hr post initiation of glucose feed. The OD₆₀₀ was 35 at time of induction, then 48.5, 61.5, 67 at 1-3 hrs post induction. Viable colony counts decreased from 0-3 hr induction [21 (13), 0, 0, 0, dilution 3 utilized 3 µl of dilution 2 cells] with numbers in parenthesis for the indicating microcolonies. Of 28 colonies scored at the time of induction, 23 retained the pHisBotA(syn)kan lacIq T7lac plasmid (kan resistant), 22 contained the chaperone plasmid (chloramphenicol resistant) and no colonies at induction grew on IPTG+Kan plates (no mutations detected). These results were indicative of very strong promoter induction, since colony viability dropped immediately after induction.

Total and soluble extracts were resolved on a 12.5% SDS-PAGE gel and stained with Coomassie. High level induction of Gro chaperones was observed, but very low level expression of soluble BotA protein was observed, increasing from 1 to 4.0 hrs post induction (no expression detected in uninduced cells). The dramatically lower expression of the BotA antigen in the presence of chaperone may be due to promoter occlusion (*i.e.*, the stronger T7 promoter on the chaperone plasmid is preferentially utilized).

b) Fermentation Of pHisBotA(syn)kan lacIq <u>T7/</u> pACYCGro BL21(DE3) Cells

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A fermentation utilizing the T7-driven BotA expression plasmid was performed. Induction was with 1 gm IPTG at 2 hrs post initiation of glucose feed. The OD₆₀₀ was 41 at time of induction, then 51.5, 61.5, 61.5 and 66 at 1-4 hrs post induction. Viable colony counts decreased from 0-4 hrs induction [71, 1 (34), 1 (1), 1, 0; dilution 3 utilized 6 μl dilution 2 cells) with numbers in parenthesis for the uninduced timepoint indicating microcolonies. Of 65 colonies scored at the time of induction, all 65 retained both the pHisBotA(syn)kan lacIq T7 plasmid (kan resistant) and the chaperone plasmid (chloramphenicol resistant) and no colonies at induction grew on IPTG+Kan plates (no mutations detected).

Total and soluble extracts were resolved on a 12.5% SDS-PAGE gel and stained with Coomassie. High level induction of Gro chaperones and moderate level expression of soluble BotA protein was observed, increasing from 1 to 4.0 hrs post induction (no expression detected in uninduced cells).

A PEI-clarified lysate (0.2% final cocnentration PEI) [850 ml from 130 gm cell pellet (2 liters fermentation harvest)] was purified on a large scale IDA column. A total of 78 mg of protein was eluted. Extracts from the purification were resolved on a 12.5% SDS-PAGE gel and stained with Coomassie. The elution was found to contain an approximately 1:1 mix of BotA/chaperone protein (Figure 32). PEI lysates prepared in this manner were typically 16 OD₂₈₀/ml. This was estimated to be 8 mg protein/ml of lysate (by BCA assay). Thus, the eluted recombinant BotA protein represented 0.55% of the total soluble cellular protein applied to the column.

In Figure 32, lane 1 contains molecular weight markers, lanes 2-9 contain extracts from pHisBotA(syn)kan lacIq T7/pACYCGro/BL21(DE3) cells before or during purification on the IDA column. Lane 2 contains total protein extract; lane 3 contains soluble protein extract; lanes 4 and 5 contain PEI-clarified lysates (duplicates); lanes 6 and 7 contain flow-through from the IDA column (duplicates) and lanes 8 and 9 contain IDA column elute (lane 9 contains 1/10 the amount applied to lane 8).

These results demonstrate, that although the majority of the BotA protein produced was insoluble, 20 mg/liter of soluble recombinant BotA protein can be purified utilizing the pHisBotA(syn)kan lacIq T7/pACYCGto/BL21(DE3) expression system.

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EXAMPLE 34

Purification Of Recombinant BotA Protein From Folding Chaperones

In this example of size exclusion chromatography was used to purify the recombinant BotA protein away from the folding chaperones and imidazole present in the IDA-purified material (Ex. 33).

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To enhance the solubility of the recombinant BotA protein during scale-up, the protein was co-expressed with folding chaperones (Ex. 33). As observed with the recombinant BotB protein (Example 40 below), the folding chaperones co-eluted with the recombinant BotA protein during the Ni-IDA purification step. Because the recombinant BotA and BotB proteins have similar molecular weights (about 1/10 the size of the non-reduced folding chaperone) and the imidazole step gradient strategy was unsuccessful in purifying BotB away from the folding chaperone (see Ex. 40), size exclusion chromatography was examined for the ability to purify the recombinant BotA protein away from the folding chaperones.

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A column (2.5 x 24 cm) containing Sephacryl S-100 HR (Pharmacia) was poured (bed volume ~ 110 ml). Proteins having molecular weights greater than 100 K are expected to elute in the void volume under these conditions and smaller proteins should be retained by the beads and elute at different times, depending on their molecular weights. To maintain solubility of the purified BotA protein, the Sephacryl column was equilibrated in a buffer having the same salt concentration as the buffer used to elute the BotA protein from the IDA column (i.e., 50 mM sodium phosphate, 0.5 M NaCl, 10% glycerol; all reagents from Mallinkrodt, Chesterfield, MO).

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Five milliliters of the IDA-purified recombinant BotA protein (Ex. 33) was filtered through a 0.45 μ syringe filter, applied to the column and the equilibration

buffer was pumped through the column at a flow rate of 1 ml/minute. Eluted proteins were monitored by absorbance at 280 nm and collected either manually or with a fraction collector (BioRad). Appropriate fractions were pooled, if necessary, and the protein was quantitated by absorbance at 280 nm and/or BCA protein assay (Pierce). The isolated peaks were then analyzed by native and/or SDS-PAGE to identify the proteins present and to evaluate purity. The folding chaperone eluted first, followed by the recombinant BotA protein and then the imidazole peak.

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SDS-PAGE analysis (12.5% polyacrylamide, reduced samples) was used to evaluate the purity of the IDA-purified recombinant BotA protein before and after S-100 purification. Figure 33 shows the difference in purity before and after the S-100 purification step. In Figure 33, lane 1 contains molecular weight markers (BioRad broad range). Lane 2 shows the IDA-purified recombinant BotA protein preparation, which is contaminated with significant amounts of the folding chaperone. Following S-100 purification, the amount of folding chaperone present in the BotA sample is reduced dramatically (lane 3). Lane 4 contains no protein (i.e., it is a blank lane); lanes 5-8 contain samples of IDA-purified recombinant BotB and BotE proteins and are discussed infra.

Endotoxin levels in the S-100 purified BotA preparation were determined using the LAL assay (Associates of Cape Cod) as describe in Example 24. The purified BotA preparation was found to contain 22.7 to 45.5 EU/mg recombinant protein.

These results demonstrate that size exclusion chromatography was successful in purifying the recombinant BotA protein from folding chaperones and imidazole following an initial IDA purification step. Furthermore, these results demonstrate that the S-100 purified BotA protein was substantially free of endotoxin.

EXAMPLE 35

Cloning And Expression Of The C Fragment
Of The C. botulinum Serotype B Toxin Gene

The C. botulinum type B neurotoxin gene has been cloned and sequenced [Whelan et al. (1992) Appl. Environ. Microbiol. 58:2345 and Hutson et al. (1994) Curr. Microbiol. 28:101]. The nucleotide sequence of the toxin gene derived from the Eklund 17B strain (ATCC 25765) is available from the EMBL/GenBank sequence data banks under the accession number X71343; the nucleotide sequence of the coding region is listed in SEQ ID NO:39. The amino acid sequence of the C. botulinum type B neurotoxin derived from the strain Eklund 17B is listed in SEQ ID NO:40. The nucleotide sequence of the C. botulinum serotype B toxin gene derived from the Danish strain is listed in SEQ ID NO:41 and the corresponding amino acid sequence is listed in SEQ ID NO:42.

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The DNA sequence encoding the native *C. botulinum* serotype B C fragment gene derived from the Eklund 17B strain can be expressed using the pETHisb vector; the resulting coding region is listed in SEQ ID NO:43 and the corresponding amino acid sequence is listed in SEQ ID NO:44. The DNA sequence encoding the native *C. botulinum* serotype B C fragment gene derived from the Danish strain can be expressed using the pETHisb vector; the resulting coding region is listed in SEQ ID NO:45 and the corresponding amino acid sequence is listed in SEQ ID NO:46. The C fragment region from any strain of *C. botulinum* serotype B can be amplified and expressed using the approach illustrated below using the C fragment derived from *C. botulinum* type B 2017 strain.

The C botulinum type B neurotoxin gene is synthesized as a single polypeptide chain which is processed to form a dimer composed of a light and a heavy chain linked via disulfide bonds; the type B neurotoxin has been reported to exist as a mixture of predominatly single chain with some double chain (Whelan et al., supra). The 50 kD carboxy-terminal portion of the heavy chain is referred to as the C

fragment or the H_C domain. Expression of the C fragment of C. botulinum type B toxin in heterologous hosts (e.g., E. coli) has not been previously reported.

The native C fragment of the C. botulinum serotype B toxin gene was cloned and expression constructs were made to facilitate protein expression in E. coli. This example involved PCR amplification of the gene, cloning, and construction of expression vectors.

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The C fragment of the C. botulinum serotype B (BotB) toxin gene was cloned using the protocols and conditions described in Example 28 for the isolation of the native BotA gene. The C. botulinum type B 2017 strain was obtained from the American Type Culture Collection (ATCC #17843). The following primer pair was used to amplify the BotB gene: 5'-CGCCATGGCTGATACAATACTAATAGAA ATG-3' [5' primer, engineered NcoI site underlined (SEQ ID NO:47)] and 5'-GCAAG CTTTTATTCAGTCCACCCTTCATC-3' [3' primer, engineered HindIII site underlined, native gene termination codon italicized (SEQ ID NO:48)]. After cloning into the pCRscript vector, the NheI(filled)/HindIII fragment was cloned into pETHisb vector as described for BotA C fragment gene in Example 28. The resulting construct was termed pHisBotB.

pHisBotB expresses the BotB gene sequences under the transcriptional control of the T7 lac promoter and the resulting protein contains an N-terminal 10XHis-tag affinity tag. The pHisBotB expression construct was transformed into BL21(DE3) pLysS competent cells and 1 liter cultures were grown, induced and his-tagged proteins were purified utilizing a NiNTA resin (eluted in low pH elution buffer) as described in Example 28. Total, soluble and purified proteins were resolved by SDS-PAGE and detected by Coomassie staining and Western blot hybridization utilizing a chicken anti-C. botulinum serotype B toxoid primary antibody (generated by immunization of hens using C. botulinum serotype B toxoid as described in Example 3). Samples of BotA and BotE C fragment proteins were included on the gels for MW and immunogenicity comparisons. Strong immunoreactivity to only the BotB protein was detected with the anti-C. botulinum serotype B toxoid antibodies. The

recombinant BotB protein was expressed at low levels (3 mg/liter) as a soluble protein. The purified BotB protein migrated as a single band of the predicted MW (i.e., ~50kD).

These results demonstrate the cloning of the native C. botulinum serotype B C fragment gene, the expression and purification of the recombinant BotB protein as a soluble his-tagged protein in E. coli.

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EXAMPLE 36

Generation Of Neutralizing Antibodies Using The Recombinant pHisBotB Protein

The ability of the purified pHisBot protein to generate neutralizing antibodies was examined. Nine BALBc mice were immunized with BotB protein (purified as described in Ex. 35) using Gerbu GMDP adjuvant (CC Biotech). The low pH elution was mixed with Gerbu adjuvant and used to immunize mice. Each mouse received a subcutaneous injection of 100 µl antigen/adjuvant mix (15 µg antigen + 1 µg adjuvant) on day 0. Mice were subcutaneously boosted as above on day 14 and bled on day 28. Mice were subsequently boosted 1-2 weeks after bleeding and were then bled on day 70.

Anti-C. botulinum serotype B toxoid titers were determined in day 28 serum from individual mice from each group using the ELISA protocol outlined in Example 29 with the exception that the plates were coated with C. botulinum serotype B toxoid, and the primary antibody was a chicken anti-C. botulinum serotype B toxoid.

Seroconversion [relative to control mice immunized with pHisBotE antigen (described below)] was observed with all 9 mice immunized with the purified pHisBotB protein.

The ability of the anti-BotB antibodies to neutralize native C. botulinum type B toxin was tested in a mouse-C. botulinum neutralization model using pooled mouse serum (see Ex. 23b). The LD₅₀ of purified C. botulinum type B toxin complex (Dr. Eric Johnson, University of Wisconsin, Madison) was determined by a intraperitoneal (IP) method [Schantz and Kautler (1978), supra] using 18-22 g female ICR mice. The amount of neutralizing antibodies present in the serum of the immunized mice was

determined using serum antibody titrations. The various serum dilutions (0.01 ml) were mixed with 5 LD₅₀ units of *C. botulinum* type B toxin and the mixtures were injected IP into mice. The neutralizations were performed in duplicate. The mice were then observed for signs of botulism for 4 days. Undiluted serum (day 28 or day 70) was found to protect 100% of the injected mice while the 1:10 diluted serum did not. This corresponds to a neutralization titer of 0.05-0.5 IU/ml.

These results demonstrate that seroconversion occurred and neutralizing antibodies were induced when the pHisBotB protein was utilized as the immunogen.

EXAMPLE 37

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Construction Of Vectors To Facilitate Expression
Of His-Tagged BotB Protein In Fermentation Cultures

A number of expression vectors were constructed to facilitate the expression of recombinant BotB protein in large scale fermentation culture. These constructs varied as to the strength of the promoter utilized (T7 or T7lac) and the presence of repressor elements (lacIq) on the plasmid. The resulting constructs varied in the level of expression achieved and in plasmid stability which facilitated the selection of a optimal expression system for fermentation scaleup.

The BotB expression vectors created for fermentation culture were engineered to utilize the kanamycin rather than the ampicillin resistance gene, and contained either the T7 or T7lac promoter, with or without the lacIq gene for the reasons outlined in Example 30.

In all cases, the protein expressed by the various expression vectors is the pHisBot B protein described in Example 35, with the only differences between clones being the alteration of various regulatory elements. Using the designations outlined below, the pHisBotB clone (Ex. 35) is equivalent to pHisBotB amp T7lac.

a) Construction Of pHisBotB kan T7lac

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pHisBotB kan T7lac was constructed by insertion of the Bg/II/HindIII fragment of pHisBotB which contains the BotB gene sequences into the pPA1870-2680 kan T7lac vector which had been digested with Bg/II and HindIII (the pPA1870-2680 kan T7lac vector contains the pET24 kan gene in the pET23 vector, such that no lacIq gene is present). Proper construction of pHisBotB kan T7lac was confirmed by restriction digestion.

b) Construction Of pHisBotB kan lacIq T7lac

pHisBotB kan lacIq T7lac was constructed by insertion of the *BglII/HindIII* fragment of pHisBotB which contains the BotB gene sequences into similarly cut pET24a vector. Proper construction of pHisBotB kan lacIq T7lac was confirmed by restriction digestion.

c) Construction Of pHisBotB kan lacIq T7

pHisBotB kan lacIq T7 was constructed by inserting the Ndel/XhoI fragment from pHisBotE kan lacIq T7lac which contains the BotB gene sequences into similarly cleaved pPA1870-2680 kan lacIq T7 vector (this vector contains the T7 promoter, the same N-terminal his-tag as the Bot constructs, the C. difficile toxin A insert, and the kan lacIq genes; this cloning replaces the C. difficile toxin A insert with the BotB insert). Proper construction was confirmed by restriction digestion.

Expression of recombinant BotB protein from these expression vectors and purification of the BotB protein is described in Example 38 below.

EXAMPLE 38

Fermentation And Purification Of Recombinant BotB Protein Utilizing The pHisBotB kan lacIq T7lac, pHisBotB kan T7lac And pHisBotB kan lacIq T7 Vectors

The pHisBotB kan lacIq T7lac, pHisBotB kan T7lac and BotB kan lacIq T7 constructs [all transformed into the Bl21(DE3) strain] were grown in fermentation cultures to determine the utility of the various constructs for large scale expression and purification of soluble BotB protein. All fermentations were performed as described in Example 31.

a) Fermentation Of pHisBotB kan lacIq T7lac/Bl21(DE3)

Cells

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The fermentation culture was induced 45 min post start of glucose feed with 1 gm IPTG (final concentration = 0.4 mM). pH was maintained at 6.5 rather than 7.0. The OD₆₀₀ was 27 at time of induction, then 35, 38, and 40 at 1-3 hrs post induction. Duplicate platings of diluted 1 hr induction samples (dilutions were prepared as described Ex. 31, dilution 3 utilized 3 μ l of dilution 2 cells) on TSA and LB+kan plates yielded 89 TSA colonies and 81 kan colonies (90% kan resistant).

Total and soluble protein extracts were resolved on a 12.5% SDS-PAGE gel and total protein was detected by staining with Coomassie blue. Low level induction of insoluble Bot B protein was observed, increasing from 1 to 3 hrs post induction (no expression was detected in uninduced cells).

b) Fermentation Of pHisBotB kan T7lac/Bl21(DE3) Cells

The fermentation culture was induced 1 hr post start of glucose feed with 2 gm IPTG (final concentration = 0.8 mM). pH was maintained at 6.5 rather than 7.0. The OD₆₀₀ was 24.5 at time of induction, then 31.5, 32, and 33 at 1-3 hrs post induction, respectively. Duplicate platings of diluted 0 hr and 2 hr induction samples (dilutions were prepared as described Ex. 31; dilution 3 utilized 3 μ l of dilution 2 cells) on TSA and LB+kan plates yielded 32 TSA colonies and 54 kan colonies (all kan resistant) for

uninduced cells, and 1 TSA colony and 0 kan colonies 2 hr post induction. These results were indicative of strong induction, since viable counts decreased dramatically 2 hrs post induction.

Total and soluble extracts were resolved on a 10% SDS-PAGE gel and total protein was detected by staining with Coomassie blue. Moderate induction of insoluble BotB protein was observed, increasing from 1 to 3 hrs post induction (no expression was detected in uninduced cells).

c) Fermentation Of pHisBotB kan lacIq T7/Bl21(DE3) Cells

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The fermentation was induced 2 hr post start of glucose feed with 4 gm IPTG (final concentration = 1.6 mM). pH was maintained at 6.5 rather than 7.0. The OD₆₀₀ was 45 at time of induction, then 47, 50, and 50 and 55 at 1-4 hrs post induction, respectively. Viable colony counts decreased after induction (96, 1, 1, 2, 3; dilution 3 utilized 3 μ l of dilution 2 cells). Of 63 colonies scored at the time of induction, all 63 retaining the BotB plasmid (kan resistant) and no colonies at induction grew on IPTG + Kan plates (no mutations detected).

Total and soluble extracts were resolved on a 12.5% SDS-PAGE gel and total protein was detected by staining with Coomassie blue. Moderate level induction of insoluble BotB protein was observed, increasing from 1 to 4 hrs post induction (lower level expression was detected in uninduced cells, since the T7 rather than T7lac promoter was utilized).

d) Purification Of pHisBotB Protein From pHisBotB amp T7lac/Bl21(DE3) Cells

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Soluble recombinant BotB protein was purified utilizing NiNTA resin from 80 ml of cell lysate generated from cells harvested from a pHisBotB fermentation [using the pHisBotB amp T7lac/BI21(DE3) strain]. As predicted from the small scale results above, the majority of the induced protein was insoluble. As well, the eluted material was contaminated with multiple *E. coli* contaminant proteins. A Coomassie blue-

stained SDS-PAGE gel containing extracts derived from pHisBotB amp T7lac/Bl21(DE3) cells before and during purification is shown in Figure 34. In Figure 34, lane 1 contains broad range protein MW markers (BioRad). Lanes 2-5 contain extracts prepared from pHisBotB amp T7lac/Bl21(DE3) cells grown in fermentation culture; lane 2 contains total protein; lane 3 contains soluble protein; lane 4 contains protein which did not bind to the NiNTA column (i.e., the flow-through) and lane 5 contains protein eluted from the NiNTA column.

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Similar results were obtained using a small scale IDA column utilizing a cell lysate from the pHisBotB kan lacIq T7 fermentation described above. 250 mls of a 20% w/v PEI clarified lysate (50 gms cell pellet) of botB kan lacIq T7/Bl21(DE3) cells were purified on a small scale IDA column. The total yield of eluted protein was 21 mg protein (assuming 1 mg/ml solution = 2 OD₂₈₀/ml). When analyzed by SDS-PAGE and Coomassie staining, the BotB protein was found to comprise approximately 50% of the eluted protein with the remainder being a ladder of *E. coli* proteins similar to that observed with the NiNTA purification.

The NiNTA alkaline phosphatase conjugate was utilized to detect his-tagged proteins on a Western blot containing total, soluble, soluble (PEI clarified), soluble (after IDA column) and elution samples from the IDA column purification. The results demonstrated that a small percentage of BotB protein was soluble, that the soluble protein was not precipitated by PEI treatment and was quantitatively bound by the IDA column. Since a l liter fermentation harvest yielded a 67.5 gm cell pellet, this indicated that the yield of soluble affinity purified BotB protein from the IDA column was 14 mg/liter.

EXAMPLE 39

Co-Expression Of Recombinant BotB Proteins And Folding Chaperones In Fermentation Cultures

Fermentations were performed to determine if the simultaneous overexpression of folding chaperones (i.e., the Gro operon) and the BotB protein resulted in enhanced solubility of the Bot B protein. This example involved fermentation of the pHisBotBkan lacIq T7lac/pACYCGro BL21(DE3), pHisBotB kan T7lac/pACYCGro Bl21(DE3) and pHisBotBkan lacIq T7/ pACYCGro BL21(DE3) cell lines. Fermentation was carried out as described in Example 31; 34 µg/ml chloramphenicol was included in the feeder and fermentation cultures.

a) Fermentation Of pHisBotBkan lacIq T7lac/pACYCGro BL21(DE3) Cells

Induction was with 4 gms IPTG at 1 hr 15 min post initiation of the glucose feed. The OD₆₀₀ was 38 at time of induction, then 50, 58.5, 62 and 68 at 1-4 hrs post induction. Viable colony counts decreased during induction (24, 0, 0, 2, 0 at 0-4 hr induction; dilution 3 utilized 3 µl of dilution 2 cells). Of 24 colonies scored at the time of induction, 24 retained the BotB plasmid (kan resistant), 24 contained the chaperone plasmid (chloramphenicol resistant) and no colonies at induction grew on IPTG+Kan plates (no mutations detected).

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Total and soluble extracts were resolved on 12.5% SDS-PAGE gels and were either stained with Coomassie blue or subjected to Western blotting (his-tagged proteins were detected utilizing the NiNTA-alkaline phosphatase conjugate). This analysis revealed that the Gro chaperones were induced to high levels, but very low level expression of soluble BotB protein was observed, increasing from 1 to 4.0 hrs post induction (no expression detected in uninduced cells, induced protein detected only on Western blot). The dramatically lower expression of BotB protein in the presence of chaperone may be due to promoter occlusion (*i.e.*, the stronger T7 promoter on the chaperone plasmid was preferentially utilized).

b) Fermentation Of pHisBotB kan T7lac/pACYCGro/Bl21(DE3) Cells

Induction was with 4 gms IPTG at 1 hr post initiation of the glucose feed. The OD₆₀₀ was 33.5 at time of induction, then 44, 51, 58.5 and 69 at 1-4 hrs post induction. Viable colony counts decreased after 2 hrs induction (43, 65, 74, 0 (70), 0 (70) at 0-4 hr induction; bracketed numbers represent microcolonies; dilution 3 utilized 3 µl of dilution 2 cells). Most colonies at induction retained the BotB plasmid (kan resistant) and the chaperone plasmid (chloramphenicol resistant) and no colonies at induction grew on IPTG+Kan plates (no mutations detected).

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Total and soluble extracts were resolved on a 12.5% SDS-PAGE gel and subjected to Western blotting; his-tagged proteins were detected utilizing the NiNTA-alkaline phosphatase conjugate. This analysis revealed that the Gro chaperones were induced to high levels and low level expression of soluble Bot B protein was observed, increasing from 1 to 4.0 hrs post induction (no expression detected in uninduced cells).

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A small scale IDA purification of BotB protein from a 250 ml PEI clarified 15% w/v extract (37.5 gm cell pellet) yielded approximately 12.5 mg protein, of which approximately 50% was BotB protein and 50% was GroEL chaperone (assessed by Coomassie staining of a 10% SDS-PAGE gel). The NiNTA alkaline phosphatase conjugate was utilized to detect his-tagged proteins on a Western blot containing total, soluble, soluble (PEI clarified), soluble (after IDA column) and elution samples from the IDA column purification. The results demonstrated that all of the BotB protein produced by the pHisBotB kan T7lac/pACYCGro/Bl21(DE3) cells was soluble; the BotB protein was not precipitated by PEI treatment and was quantitatively bound by the IDA column. Since a 1 liter fermentation harvest yielded a 75 gm cell pellet, this indicated that the yield of soluble affinity purified bot B protein from this fermentation was 12.5 mg/liter. These results also demonstrated that additional purification steps are necessary to separate the chaperone proteins from the BotB protein.

c) Fermentation Of pHisBotBkan lacIq T7/pACYCGro/BL21(DE3) Cells

Induction was with 4 gms IPTG at 2 hr post initiation of the glucose feed. The OD₆₀₀ was 46 at time of induction, then 56, 63, 69 and 71.5 at 1-4 hrs post induction. Viable colony counts decreased after induction (58, 3(5), 3, 0, 0 at 0-4 hr induction; bracketed numbers represent microcolonies; dilution 3 utilized 3 µl of dilution 2 cells). All (53/53) colonies scored at the time of induction retained the BotB plasmid (kan resistant) and the chaperone plasmid (chloramphenicol resistant) and no colonies at induction grew on IPTG+Kan plates (no mutations detected).

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Total and soluble extracts were resolved on a 10% SDS-PAGE gels and Western blotted and his-tagged proteins were detected utilizing the NiNTA-alkaline phosphatase conjugate. This analysis revealed that the Gro chaperones were induced to high levels (observed by ponceau S staining), and a much higher expression of soluble Bot B protein (compared to expression in the pHisBotB kan T7lac/pACYCGro fermentation) was observed at all timepoints, including uninduced cells (some increase in BotB protein levels were observed after induction).

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A small scale IDA purification of BotB protein from a 100 ml PEI clarified 15% w/v extract (15 gm cell pellet) yielded approximately 40 mg protein, of which approximately 50% was BotB protein and 50% was GroEL chaperone, as assessed by Coomassie staining of a 10% SDS-PAGE gel. The NiNTA alkaline phosphatase conjugate was utilized to detect his-tagged proteins on a Western blot containing total, soluble, soluble (PEI clarified), soluble (after IDA column) and elution samples from the IDA column purification. The results demonstrated that a significant percentage (i.e., ~10-20 %) of BotB protein was soluble, that the solubilized protein was not precipitated by PEI treatment and was quantitatively bound by the IDA column. Since a 10 liter fermentation yielded a 108 gm cell pellet, this indicated that the yield of soluble affinity purified BotB protein from this fermentation was 144 mg/liter.

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In a scale up experiment, 2 liters of a 20% w/v PEI clarified lysate of pHisBotB kan laclq T7/pACYCGro/BL21(DE3) cells were purified on a large scale IDA column. The purification was performed in duplicate. The total yield of BotB

protein was 220 and 325 mgs protein in the two experiments (assuming 1 mg/ml solution = 2.0 OD₂₈₀/ml). This represents 0.7% or 1.0%, respectively, of the total soluble cellular protein (assuming a PEI lystate having a concentration of 8 mg protein/ml and that the eluted material comprises a 1:1 mixture of BotB and folding chaperone). The NiNTA alkaline phosphatase conjugate was utilized to detect histagged proteins on a Western blot containing total, soluble, soluble (PEI clarified), soluble (after IDA column) and elution samples from the IDA column purification. These results demonstrated that a significant percentage (i.e., ~10-20 %) of the BotB protein was soluble, that the solubilized protein was not precipitated by PEI treatment and was quantitatively bound by the IDA column. Since a 1 liter fermentation harvest yielded a 108 gm cell pellet, this indicated that the yield of soluble affinity purified BotB protein from the large scale purification was 60 mg or 89 mg/liter. These results also demonstrated that further purification would be necessary to remove the contaminating chaperone protein.

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The above results provide methodologies for the purification of soluble BotB protein from fermentation cultures, in a form contaminated predominantly with a single *E. coli* protein (the folding chaperone utilized to enhance solubility). In the next example, methods are provided for the removal of the contaminating chaperone protein.

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EXAMPLE 40

Removal Of Contaminating Folding Chaperone Protein

From Purified Recombinant C. botulinum Type B Protein

In this example size exclusion chromatography and ultrafiltration was used to purify recombinant BotB protein from the folding chaperones and imidazole in IDA-purified material.

To enhance the solubility of the recombinant BotB protein during scale-up, the protein was co-expressed with folding chaperones (see Ex. 39). During the Ni-IDA purification step, the folding chaperones co-eluted with the BotB protein in 800 mM

imidazole; therefore, a second purification step was required to isolate the BotB free of folding chaperones. Lane 3 of Figure 35 contains proteins eluted from an IDA column to which a lysate of pHisBotB kan lacIq T7/pACYCGro/BL21(DE3) cells had been applied; the proteins were resolved on a 4-15% polyacrylamide pre-cast gradient gel (Bio-Rad, Hercules, CA) run under native conditions and then stained with Coomassie blue. In Figure 35, lanes 1 and 4 contain proteins present in peak 1 and peak 2 from a Sephacryl S-100 column run as described below; lane 2 is blank.

As seen in lane 3 of Figure 35, the IDA-purified sample consists primarily of the folding chaperones and the BotB protein. The fact that the chaperones and the BotB antigen appear as two distinct bands under native conditions suggested they were not complexed together and therefore, it should be possible to separate them, using either a gradient of imidazole concentrations or size exclusion methods.

In order to determine whether a gradient of imidazole concentrations could be used to separate the chaperone from the BotB protein, a step gradient using imidazole at 200, 400, 600, and 800 mM in 50 mM sodium phosphate, 0.5 M NaCl and 10 % glycerol, pH 6.8 was applied to an IDA column (containing proteins bound from a lysate of pHisBotB kan lacIq T7/pACYCGro/BL21(DE3) cells). By narrowing the range of imidazole concentrations, it was hoped that the BotB and chaperone proteins would differentially elute at different concentrations of imidazole. Eluted proteins were monitored by absorbance at 280 nm and collected either manually or with a fraction collector (BioRad). Protein was found to elute at 200 and 400 mM imidazole only.

Figure 36 shows a Coomassie stained SDS-PAGE gel containing protein eluted during the imidazole step gradient. Lane 1 contains broad range MW markers (BioRad). Lane 2 contains BotB protein purified by IDA chromatography of an extract of pHisBotB/BL21(DE3) pLysS cells grown in shaker flask culture (i.e., no coexpression of chaperones; Ex. 35). Lane 3 contains a 20% w/v PEI clarified lysate of pHisBotB kan lacIq T7/pACYCGro/BL21(DE3) cells (i.e., the lysate prior to purification by IDA chromatography). Lanes 4 and 5 contain protein which eluted at

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200 or 400 mM imidazole, respectively. Lane 6 is blank. Lanes 7 and 8 contain 1/5 the load present in lanes 4 and 5.

As shown in Figure 36, both the chaperone and the BotB protein eluted in 200 mM imidazole, and more chaperone elutes in 400 mM imidazole, however no concentration of imidazole tested permitted the elution of BotB protein alone. Consequently, no significant purification was achieved using imidazole at these concentrations.

Because of the considerable difference in molecular weights between the folding chaperone, which is a multimer with a total molecular weight around 400 kD (as determined on a Shodex KB 804 sizing column by HPLC), and the recombinant BotB protein (molecular weight around 50 kD), size exclusion chromatography was next examined for the ability to separate these proteins.

a) Size Exclusion Chromatography

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A column containing Sephacryl S-100 HR (S-100) (Pharmacia) was poured (2.5 cm x 24 cm; ~110 ml bed volume). The column was equilibrated in a buffer consisting of phosphate buffered saline (10mM potassium phosphate, 150 mM NaCl, pH 7.2) and 10 % glycerol (Mallinkrodt). Typically, 5 ml of the IDA-purified BotB protein was filtered through a 0.45 μ syringe filter and applied to the column, and the equilibration buffer was pumped through the column at a flow rate of 1 ml/minute. Eluted proteins were monitored by absorbance at 280 nm and collected either manually or with a fraction collector. Appropriate tubes were pooled, if necessary, and the protein was quantitated by absorbance at 280 nm and/or by BCA protein assay. The isolated peaks were then analyzed by native and/or SDS-PAGE to identify the protein and evaluate the purity.

Because of its larger size, the folding chaperone eluted first, followed by the recombinant BotB protein. A smaller third peak was observed which failed to stain when analyzed by SDS-PAGE and therefore was presumed to be imidazole.

SDS-PAGE analysis (12.5% polyacrylamide, reduced samples) was used to evaluate the purity of the IDA-purified recombinant BotB protein before and after S-100 purification. The results are shown in Figure 33.

In Figure 33, lane 1 contains broad range MW markers (BioRad). Lane 5 contains IDA-purified BotB protein. Lane 6 contains IDA-purified BotB protein following S-100 purification. Lane 7 is blank (lanes 2-4 were discussed in Ex. 34 above).

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The results shown in Figure 33 show that the IDA-purified BotB is significantly contaminated with the folding chaperone (molecular weight about 60 kD under reducing conditions; lane 6). Following S-100 purification, the amount of folding chaperone present in the BotB sample was reduced dramatically (lane 7). Visual inspection of the Coomassie stained SDS-PAGE gel revealed that after S-100 purification, > 90% of the total protein present was BotB.

The IDA-purified BotB and the S-100-purified BotB samples were analyzed by HPLC on a size exclusion column (Shodex KB 804); this analysis revealed that the BotB protein represented 64% of the total protein in the IDA-purified sample and that following S-100 purification, the BotB protein represented >95% of the total protein in the sample.

The IDA-purified BotB material was also applied to a ACA 44 (SpectraPor, Houston, TX) column. The ACA 44 resin is equivalent to the S-100 resin and chromatography using the ACA 44 resin was carried out exactly as described above for the S-100 resin. The ACA 44 resin was found to separate the recombinant BotB protein from the folding chaperone. The ACA 44-purified BotB sample was analyzed for endotoxin using the LAL assay (Associates of Cape Cod) as describe in Example 24. Two aliquouts of the ACA 44-purified BotB preparation were analyzed and were found to contain either 58 to 116 EU/mg recombinant protein or 94 to 189 EU/mg recombinant protein.

These results demonstrate that size exclusion chromatography can be used to purify the recombinant BotB protein from the folding chaperone and imidazole in IDA-purified material.

b) Ultrafiltration For The Separation Of Recombinant BotB Protein And Chaperones

Ultrafiltration was examined as an alternative method for the separation recombinant BotB protein and folding chaperones in IDA-purified material. While in this example only mixtures of BotB and chaperones were separated by ultrafiltration, this technique is suitable for use with recombinant BotA and BotE proteins as well provided that the wash buffers used are altered as necessary to take into account different requirements for solubility.

The recombinant BotB protein and folding chaperones were separated using a two-step sequential ultrafiltration method. The first membrane used had a nominal molecular weight cutoff (MWCO) of approximately 100 kD; this membrane retains the larger folding chaperone while allowing the smaller recombinant protein to pass through. The addition of several volumes of wash buffer may be required to efficiently wash the recombinant protein through the membrane. The second step utilized a membrane with a nominal MWCO of approximately 10 kD. During this step, the recombinant antigen was retained by the membrane and could be concentrated to the degree desired and the imidazole and excess wash buffer passed through the membrane.

Twenty-seven milliliters of an IDA-purified BotB preparation was ultrafiltered through a 47 mm YM 100 (100 kD MWCO) membrane (Amicon) in a 50 ml stirred cell (Amicon). The membrane was washed in dd H₂O prior to use as recommended by the manufacturer. Six volumes of 10% glycerol in PBS were washed through to remove most of the recombinant BotB protein and this wash was collected in a separate vessel. The resulting BotB protein-rich filtrate was then concentrated 12-fold using a YM 10 (10 kD MWCO) membrane (Amicon), to a final volume of 14 ml. The YM 100 and YM 10 concentrates were analyzed along with the lysate starting material by native PAGE using a 4-15% pre-cast gradient gel (BioRad). The results are shown in Figure 37.

In Figure 37, lane 1 contains IDA-purified BotB derived from a shaker flask culture (i.e., no co-expression of chaperones; Ex. 35); lane 2 contains a 20% w/v PEI

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clarified lysate of pHisBotB kan lacIq T7/pACYCGro/BL21(DE3) cells; lane 3 shows the lysate of lane 3 after IDA purification; lane 4 contains the YM 10 concentrate and lane 5 contains the YM 100 concentrate.

The results shown in Figure 37 demonstrate that the recombinant BotB protein can be purified away from the folding chaperone by ultrafiltration through a 100 kD MWCO membrane (lane 4), leaving the chaperone protein in the 100 kD concentrate (lane 5). Analysis of the sample in lane 5 also showed that very little of the BotB protein was retained by the 100 kD MWCO membrane after 6 volumes of wash buffer had been applied.

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The BotB samples following IDA chromatography and following ultrafiltration through the YM 100 membrane were anlyzed by HPLC on a size exclusion column (Shodex KB 804); this analysis revealed that the BotB protein represented 64% of the total protein in the IDA-purified sample and that following ultrafiltration through the YM 100 membrane, the BotB protein represented >96% of the total protein in the sample.

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The BotB protein purified by ultrafiltration through the YM 100 membrane was examined for endotoxin using the LAL assay (Associates of Cape Cod) as describe in Example 24. Two aliquouts of the YM 100-purified BotB preparation were analyzed and were found to contain either 18 to 36 EU/mg recombinant protein or 125 to 250 EU/mg recombinant protein.

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The above results demonstrate that size exclusion chromatography and ultrafiltration can be used to purify recombinant botulinal toxin proteins away from folding chaperones.

EXAMPLE 41

Cloning And Expression Of The C Fragment Of The C. botulinum Serotype E Toxin Gene

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The *C. botulinum* type E neurotoxin gene has been cloned and sequenced from several different strains [Poulet *et al.* (1992) Biochem. Biophys. Res. Commun. 183:107 (strain Beluga); Whelan *et al.* (1992) Eur. J. Biochem. 204:657 (strain NCTC 11219); Fujii *et al.* (1990) Microbiol. Immunol. 34:1041 (partial sequence of strains Mashike, Iwani and Otaru) and Fujii *et al.* (1993) J. Gen. Microbiol. 139:79 (strain Mashike)]. The nucleotide sequence of the type E toxin gene is available from the EMBL sequence data bank under accession numbers X62089 (strain Beluga) and X62683 (strain NCTC 11219). The nucleotide sequence of the coding region (strain Beluga) is listed in SEQ ID NO:49. The amino acid sequence of the *C. botulinum* type E neurotoxin derived from strain Belgua is listed in SEQ ID NO:50. The nucleotide sequence of the coding region (strain NCTC 11219) is listed in SEQ ID NO:51. The amino acid sequence of the *C. botulinum* type E neurotoxin derived from strain NCTC 11219 is listed in SEQ ID NO:52.

The DNA sequence encoding the native C. botulinum serotype E C fragment gene derived from the Beluga strain can be expressed as a histidine-tagged protein using the pETHisb vector; the resulting coding region is listed in SEQ ID NO:53 and the corresponding amino acid sequence is listed in SEQ ID NO:54. The DNA sequence encoding the C fragment of the native C. botulinum serotype E gene derived from the NCTC 11219 strain can be expressed as a histidine-tagged fusion protein using the pETHisb vector; the resulting coding region is listed in SEQ ID NO:55 and the corresponding amino acid sequence is listed in SEQ ID NO:56. The C fragment region from any strain of C. botulinum serotype E can be amplified and expressed using the approach illustrated below using the C fragment derived from C. botulinum type E 2231strain (ATCC #17786).

The type E neurotoxin gene is synthesized as a single polypeptide chain which may be converted to a double-chain form (i.e., a heavy chain and a light chain) by cleavage with trypsin; unlike the type A neurotoxin, the type E neurotoxin exists essentially only in the single-chain form. The 50 kD carboxy-terminal portion of the heavy chain is referred to as the C fragment or the H_C domain. Expression of the C fragment of C. botulinum type E toxin in heterologous hosts (e.g., E. coli) has not been previously reported.

The native C fragment of the C. botulinum serotype E toxin (BotE) gene was cloned and inserted into expression vectors to facilitate expression of the recombinant BotE protein in E. coli. This example involved PCR amplification of the gene, cloning, and construction of expression vectors.

The BotE serotype gene was isolated using PCR as described for the BotA serotype gene in Example 28. The *C. botulinum* type E strain was obtained from the American Type Culture Collection (ATCC #17786; strain 2231). The following primer pair was used in the PCR amplification: 5'-CGCATGGCTCTTTCTTCTTAT ACAGATGAT-3' (5' primer, engineered *NcoI* site underlined) (SEQ ID NO:57) and 5'-GCAAGCTTTTATTTTTCTTGCCATCCATG-3' (3' primer, engineered *HindIII* site underlined, native gene termination codon italicized) (SEQ ID NO:58). The PCR product was inserted into pCRscript as described in Example 28. The resulting pCRscript BotE clone was confirmed by restriction digestion, as well as, by obtaining the sequence of approximately 300 bases located at the 5' end of the C fragment coding region using standard DNA sequencing methods. The resulting BotE sequence was identical to that of the published *C. botulinum* type E toxin sequence [Whelan *et al* (1992), *supra*].

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The Nhel(filled)/HindIII fragment from a pCRscript BotE recombinant was cloned into pETHisb vector as described for BotA C fragment in Example 28. The resulting construct was termed pHisBotE. pHisBotE expresses the BotE gene under the control of the T7 lac promoter and the resulting protein contains an N-terminal 10XHis-tag affinity tag.

The pHisBotE expression construct was transformed into BL21(DE3) pLysS competent cells and 1 liter cultures were grown, induced and his-tagged proteins were purified utilizing a NiNTA resin (eluted in low pH elution buffer) as described in Example 28. Total, soluble and purified proteins were resolved by SDS-PAGE and detected by Coomassie staining. The results are shown in Figure 38.

In Figure 38, lane 1 contains broad range MW markers (BioRad); lane 2 contains a total protein extract; lane 3 contains a soluble protein extract; lane 4 contains proteins present in the flow through from the NiNTA column (this sample was not diluted prior to loading and therefore represents a load 5X that of the load applied for the total and soluble extracts in lanes 2 and 3); lane 5 contains proteins eluted from the NiNTA column; lane 6 contains protein eluted from a NiNTA column which had been stored at -20°C for 1 year.

The pHisBotE protein was expressed at moderate levels (7 mg/liter) as a totally soluble protein. The purified protein migrated as a single band of the predicted MW.

Western blot hybridization utilizing a chicken anti-C. botulinum serotype E toxoid primary antibody (generated by immunization of hens as described in Example 3 using C. botulinum serotype E toxoid) was also performed on the total, soluble and purified BotE proteins. Samples of BotA and BotB C fragments were also included on the gels to facilitate MW and immunogenicity comparisons. Strong immunoreactivity was detected using the anti-C. botulinum type E toxoid antibody only with the BotE protein.

These results demonstrate that the native BotE gene sequences can be expressed as a soluble his-tagged protein in *E. coli* and purified by metal-chelation affinity chromatography.

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EXAMPLE 42

Generation Of Neutralizing Antibodies Using The Recombinant pHisBotE Protein

The ability of the purified pHisBotE protein to generate neutralizing antibodies was examined. Nine BALBc mice were immunized with BotE protein (purified as described in Ex. 41) using Gerbu GMDP adjuvant (CC Biotech). The low pH elution was mixed with Gerbu adjuvant and used to immunize mice. Each mouse received a subcutaneous injection of 100 µl antigen/adjuvant mix (35 µg antigen + 1 µg adjuvant) on day 0. Mice were subcutaneously boosted as above on day 14 and bled on day 28. Mice were subsequently boosted and bled on day 70.

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Anti-C. botulinum serotype E toxoid titers were determined in day 28 serum from individual mice from each group using the ELISA protocol outlined in Example 29 with the exception that the plates were coated with C. botulinum serotype E toxoid, and the primary antibody was a chicken anti-C. botulinum serotype E toxoid.

Seroconversion [relative to control mice immunized with the p6xHisBotA antigen (Ex. 29)] was observed with all 9 mice immunized with the purified pHisBotE protein.

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The ability of the anti-BotE antibodies to neutralize native *C. botulinum* type E toxin was tested in a mouse-*C. botulinum* neutralization model using pooled mouse serum (see Ex. 23b). The LD₅₀ of purified *C. botulinum* type E toxin complex (Dr. Eric Johnson, University of Wisconsin, Madison) was determined by a intraperitoneal (IP) method [Schantz and Kautler (1978), supra] using 18-22 g female ICR mice. The amount of neutralizing antibodies present in the serum of the immunized mice was determined using serum antibody titrations. The various serum dilutions (0.01 ml) were mixed with 5 LD₅₀ units of *C. botulinum* type E toxin and the mixtures were injected IP into mice. The neutralizations were performed in duplicate. The mice were then observed for signs of botulism for 4 days. Undiluted serum from day 28 did not protect, while undiluted, 1/10 diluted and 1/100 diluted day 70 serum protected (1005 of animals) while 1/1000 diluted day 70 serum did not. This corresponds to a neutralization titer of 50-500 IU/ml.

These results demonstrate that seroconversion occurred and neutralizing antibodies were induced when the recombinant BotE protein was utilized as the immunogen.

EXAMPLE 43

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Construction Of Vectors To Facilitate Expression
Of His-Tagged BotE Protein In Fermentation Cultures

A number of expression vectors were constructed to facilitate the expression of recombinant BotE protein in large scale fermentation culture. These constructs varied as to the strength of the promoter utilized (T7 or T7lac) and the presence of repressor elements (lacIq) on the plasmid. The resulting constructs varied in the level of expression achieved and in plasmid stability which facilitated the selection of a optimal expression system for fermentation scaleup. This example involved a) construction of BotE expression vectors and b) determination of expression levels in small scale cultures.

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a) Construction Of BotE Expression Vectors

The BotE expression vectors created for fermentation culture were engineered to utilize the kanamycin rather than the ampicillin resistance gene, and contained either the T7 or T7lac promoter, with or without the lacIq gene for the reasons outlined in Example 30.

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In all cases, the protein expressed by the various expression vectors is the pHisBotE protein described in Example 41, with the only differences between clones being the alteration of various regulatory elements. Using the designations outlined below, the pHisBotE clone (Ex. 41) is equivalent to pHisBotE amp T7lac.

i) Construction Of pHisBotE kan lacIq T7lac

pHisBotE kan laciq T7lac was constructed by inserting the Xbal/HindIII fragment of pHisBotE which contains the BotE gene sequences into Xbal/HindIII-cleaved pET24a vector. Proper construction was confirmed by restriction digestion.

ii) Construction Of pHisBotE kan T7

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pHisBotE kan T7 was constructed by ligating the BotE-containing Xbal/Sapl fragment of pHisBotE kan lacIqT7lac to the T7 promoter-containing Xbal/Sapl fragment of pET23a. Proper construction was confirmed by restriction digestion.

iii) Construction Of pHisBotE kan lacIqT7

pHisBotE kan lacIqT7 was constructed by inserting the Bg/II/HindIII fragment from pHisBotE kan T7 which contains the BotE gene sequences into Bg/II/HindIII-cleaved pET24 vector. Proper construction was confirmed by restriction digestion.

b) Determination Of BotE Expression Levels In Small Scale Cultures

The three BotE kan expression vectors described above were transformed into BI21(DE3) competent cells and 50 ml (2XYT + 40 µg/ml kan) cultures were grown and induced with ITPG as described in Example 28. Total and soluble protein extracts from before and after induction made as described in Example 28. The total and soluble extracts were resolved on a 12.5% SDS-PAGE gel, and his-tagged proteins were detected on a Western blot utilizing the NiNTA-alkaline phosphatase conjugate as described in Example 31(c)(iii). The results showed that all three BotE cell lines expressed his-tagged proteins of the predicted MW for the BotE protein upon induction. The results also demonstrated that the two constructs that contained the T7 promoter expressed the BotE protein before induction, while the T7lac promoter construct did not. Upon induction, the T7 promoter-containing constructs induced to

higher levels than the T7lac-containing construct, with the pHisBotE kan lacIqT7/Bl21(DE3) cells accumulating the maximal levels of BotE protein.

EXAMPLE 44

Expression And Purification Of pHisBotE From Fermentation Cultures

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Based on the small scale inductions performed in Example 43, the pHisBotE kan laclq T7/Bl21(DE3) strain was selected for fermentation scaleup. This example involved the fermentation and purification of recombinant BotE C fragment protein.

A fermentation with the pHisBotE kan lacIq T7/Bl21(DE3) strain was performed as described in Example 31. The fermentation culture was induced 2 hrs post start of the glucose feed with 4 gm IPTG (final concentration = 1.6 mM). The OD₆₀₀ was 42 at time of induction, then 46.5, 48, 53 and 54 at 1-4 hrs post induction. Viable colony counts decreased from 0-4 hr induction [131, 4 (28), 7 (3), 7, 8; dilution 3 utilized 6 μl of dilution 2 cells; bracketed colonies are microcolonies]. All (32/32) colonies scored at the time of induction retained the BotE plasmid (kan resistant) and no colonies at induction grew on IPTG+Kan plates (no mutations detected). These results were indicative of strong promoter induction, since colony viability reduced after induction, and the culture stopped growing during fermentation (stopped at 54 OD₆₀₀/ml).

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Total and soluble extracts were resolved on a 12.5% SDS-PAGE gel and total protein was detected by staining with Coomassie blue. The results are shown in Figure 39.

In Figure 39, lane 1 contains total protein from a pHisBotA kan T7 lac/Bl21(DE3) pLysS fermentation (Ex. 24). Lanes 2-9 contain extracts prepared from the above pHisBotE kan lacIq T7/Bl21(DE3) fermentation; lanes 2-4 contain total protein extracts prepared at 0, 1 and 2 hours post-induction, respectively. Lane 5 contains a soluble protein extract prepared at 2 hours post-induction. Lanes 6 and 7 contain total and soluble extracts prepared at 3 hours post-induction, respectively.

Lanes 8 and 9 contain total and soluble extracts prepared at 4 hours post-induction, respectively. Lane 10 contains broad range MW markers (BioRad).

The results shown in Figure 39 demonstrate that moderate level induction of totally soluble Bot E protein was observed, increasing from 1 to 4 hrs post induction (no expression was detected in uninduced cells). From a 2 liter fermentation harvest a 155 gm (wet wt) cell pellet was obtained and used to make a PEI-clarified lysate (1 liter in CRB, pH 6.8). The lysate was applied to a large scale IDA column and 200 mg of BotE protein, which was found to be greater than 95% pure (as judged by visual inspection of a Coomassie stained SDS-PAGE gel), was recovered. This represents 2.5% of the total soluble cellular protein (assuming a PEI lysate having a concentration of 8 mg protein/ml) and corresponds to a yield of 100 mg BotE protein/liter of fermentation culture.

The above results demonstrate that high levels of the recombinant BotE protein can be expressed and purified from fermentation cultures.

15 EXAMPLE 45

Removal Of Imidazole From Purified Recombinant BotE Protein Preparations

The expression of recombinant BotE protein, unlike the BotA and BotB proteins, did not require the presence of folding chaperones to maintain solubility during scale-up. A size exclusion chromatography step was included however to remove the imidazole from the sample and exchange the IDA elution buffer for one consistent with the BotA antigen.

A Sephacryl S-100 HR (S-100; Pharmacia) column was poured (2.5 cm x 24 cm; bed volume \sim 110 ml). Under these conditions, the BotE protein should be retained by the beads to a lesser degree than the smaller imidazole, therefore the BotE protein should elute from the column before the imidazole. The column was equilibrated in a buffer consisting of 50 mM sodium phosphate, 0.5 M NaCl, and 10% glycerol (all reagents from Mallinkrodt). Five milliliters of the IDA-purified BotE protein (Ex. 44) was filtered through a 0.45 μ syringe filter and applied to the S-100

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column, and equilibration buffer was pumped through the column at a flow rate of 1 ml/minute. Eluted proteins were monitored by absorbance at 280 nm, and collected either manually or with a fraction collector. Appropriate tubes were pooled if necessary, and the protein was quantitated by absorbance at 280 nm and/or BCA protein assay. The isolated peaks were then analyzed by native and/or SDS-PAGE to identify the protein(s) and evaluate the purity.

Figure 40 provides a representative chromatogram showing the purification of IDA-purified BotE on the S-100 column. Even though folding chaperones were not over-expressed with this antigen, a small amount of protein eluted at a time consistent with the folding chaperones expressed with BotA and BotB proteins (Gro) (see the first peak). The second peak in the chromatogram contained the BotE protein, and the third peak was presumably imidazole. This presumed imidazole peak was isolated in comparable levels in IDA-purified BotA and BotB protein preparations as well.

These results demonstrate that size exclusion chromatography can be used to remove imidazole and traces of contaminating high molecular weight proteins from IDA-purified BotE protein preparations.

The S-100-purified BotE protein was tested for endotoxin contamination using the LAL assay as described in Example 24. This preparation was found to contain 64 to 128 EU/mg recombinant protein and is therefore substantially free of endotoxin.

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The S-100 purified BotE was mixed with purified preparations of BotA and BotB proteins and used to immunize mice; 5 µg of each Bot protein was used per immunization and alum was included as an adjuvant. After two immunizations with this trivalent vaccine, the immunized mice were challanged with *C. botulinum* toxin. The immunized mice contained neutralizing antibodies sufficient to neutralize between 100,000 to 1,000,000 LD₅₀ of either toxin A or toxin B and between 1,000 to 10,000 LD₅₀ of toxin E. The titer of neutralizing antibodies directed against toxin E would be expected to increase following subsequent boosts with the vaccine. These results demonstrate that a trivalent vaccine containing recombinant BotA, BotB and BotE proteins provokes neutralizing antibodies.

EXAMPLE 46

Expression Of The C Fragment Of The C. botulinum

Serotype C Toxin Gene And Generation Of Neutralizing Antibodies

The C. botulinum type C1 neurotoxin gene has been cloned and sequenced [Kimura et al. (1990) Biochem. Biophys. Res. Comm. 171:1304]. The nucleotide sequence of the toxin gene derived from the C. botulinum type C strain C-Stockholm is available from the EMBL/GenBank sequence data banks under the accession number D90210; the nucleotide sequence of the coding region is listed in SEQ ID NO:59. The amino acid sequence of the C. botulinum type C1 neurotoxin derived from this strain is listed in SEQ ID NO:60.

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The DNA sequence encoding the native C. botulinum serotype C1 C fragment gene derived from the C-Stockholm strain can be expressed using the pETHisb vector; the resulting coding region is listed in SEQ ID NO:61 and the corresponding amino acid sequence is listed in SEQ ID NO:62. The C fragment region from any strain of C. botulinum serotype C can be amplified and expressed using the approach illustrated below using the C fragment derived from C. botulinum type C C-Stockholm strain. Expression of the C fragment of C. botulinum type C1 toxin in heterologous hosts (e.g., E. coli) has not been previously reported.

The C fragment of the C. botulinum serotype C1 (BotC1) toxin gene is cloned using the protocols and conditions described in Example 28 for the isolation of the native BotA gene. A number of C. botulinum serotype C strains (expressing either or both C1 and C2 toxin) are available from the ATCC [e.g., 2220 (ATCC 17782), 2239 (ATCC 17783), 2223 (ATCC 17784; a type C-β strain; C-β strains produce C2 toxin), 662 (ATCC 17849; a type C-α strain; C-α strains produce mainly C1 toxin and a small amount of C2 toxin), 2021 (ATCC 17850; a type C-α strain) and VPI 3803 (ATCC 25766)]. Alternatively, other type C strains may be employed for the isolation of sequences encoding the C fragment of C. botulinum serotype C toxin.

The following primer pair is used to amplify the BotC gene: 5'-CGCCATGGC TTTATTAAAAGATATAATTAATG-3' [5' primer, engineered Ncol site underlined (SEQ ID NO:63)] and 5'-GCAAGCTTTTATTCACTTACAGGTAC AAAACC-3' [3' primer, engineered HindIII site underlined, native gene termination codon italicized (SEQ ID NO:64)]. Following PCR amplification, the PCR product is inserted into the pCRscript vector and then the 1.5 kb fragment is cloned into pETHisb vector as described for BotA C fragment gene in Example 28. The resulting construct is termed pHisBotC. Proper construction is confirmed by DNA sequencing of the BotC sequences contained within pHisBotC.

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pHisBotC expresses the BotC gene sequences under the transcriptional control of the T7 lac promoter and the resulting protein contains an N-terminal 10XHis-tag affinity tag. The pHisBotC expression construct is transformed into BL21(DE3) pLysS competent cells and 1 liter cultures are grown, induced and his-tagged proteins are purified utilizing a NiNTA resin (eluted in 250 mM imidazole, 20% glycerol) as described in Example 28. Total, soluble and purified proteins are resolved by SDS-PAGE and detected by Coomassie staining and Western blot hybridization utilizing a Ni-NTA-alkaline phosphatase conjugate (Qiagen) which recognizes his-tagged proteins as described in Example 31(c)(iii). This analysis permits the determination of expression levels of the pHisBotC protein (i.e., number of mg/liter expressed as a soluble protein). The purified BotC protein will migrate as a single band of the predicted MW (i.e., ~50kD).

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The level of expression of the pHisBotC protein may be modified (increased) by substitution of the T7 promoter for the T7lac promoter, or by inclusion of the lacIq gene on the expression plasmid, and plasmid expressed in BL21(DE3) cell lines in fermentation cultures as described in Example 30. If only very low levels (i.e., less than 0.5%) of soluble pHisBotC protein are expressed using the above expression systems, the pHisBotC construct may be co-expressed with pACYCGro construct as described in Example 32. In this case, the recombinant BotC protein may co-purify with the folding chaperones. The contaminating chaperones may be removed as

described in Example 34. Preparations of purified pHisBotC protein are tested for endotoxin contamination using the LAL assay as described in Example 24.

The purified pHisBotC protein is used to generate neutralizing antibodies.

BALBc mice are immunized with the BotC protein using Gerbu GMDP adjuvant (CC Biotech) as described in Example 36. The ability of the anti-BotC antibodies to neutralize native C. botulinum type C toxin is demonstrated using the mouse-C. botulinum neutralization model described in Example 36.

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EXAMPLE 47

Expression Of The C Fragment Of The C. botulinum

Serotype D Toxin Gene And Generation Of Neutralizing Antibodies

The C. botulinum type D neurotoxin gene has been cloned and sequenced [Sunagawa et al. (1992) J. Vet. Med. Sci. 54:905 and Binz et al. (1990) Nucleic Acids Res. 18:5556]. The nucleotide sequence of the toxin gene derived from the CB16 strain is available from the EMBL/GenBank sequence data banks under the accession number S49407; the nucleotide sequence of the coding region is listed in SEQ ID NO:65. The amino acid sequence of the C. botulinum type D neurotoxin derived from the CB16 strain is listed in SEQ ID NO:66.

The DNA sequence encoding the native *C. botulinum* serotype D C fragment gene derived from a BotD expressing strain can be expressed using the pETHisb vector; the resulting coding region is listed in SEQ ID NO:67 and the corresponding amino acid sequence is listed in SEQ ID NO:68. The C fragment region from any strain of *C. botulinum* serotype D can be amplified and expressed using the approach illustrated below using the C fragment derived from *C. botulinum* type D CB16 strain. Expression of the C fragment of *C. botulinum* type D toxin in heterologous hosts (e.g., *E. coli*) has not been previously reported.

The C fragment of the C. botulinum serotype D (BotD) toxin gene is cloned using the protocols and conditions described in Example 28 for the isolation of the

native BotA gene. A number of C. botulinum type D strains are available from the ATCC [e.g., ATCC 9633, 2023 (ATCC 17851), and VPI 5995 (ATCC 27517)].

The following primer pair is used to amplify the BotD gene: 5'-CGCCATGGC TTTATTAAAAGATATAATTAATG-3' [5' primer, engineered Ncol site underlined (SEQ ID NO:63)] and 5'-GCAAGCTTTTACTCTACCCATCCTGGATCCCT-3' [3' primer, engineered HindIII site underlined, native gene termination codon italicized (SEQ ID NO:69)]. Following PCR amplification, the PCR product is inserted into the pCRscript vector and then the 1.5 kb fragment is cloned into pETHisb vector as described for BotA C fragment gene in Example 28. The resulting construct is termed pHisBotD.

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pHisBotD expresses the BotD gene sequences under the transcriptional control of the T7 lac promoter and the resulting protein contains an N-terminal 10XHis-tag affinity tag. The pHisBotD expression construct is transformed into BL21(DE3) pLysS competent cells and 1 liter cultures are grown, induced and his-tagged proteins are purified utilizing a NiNTA resin as described in Example 28. Total, soluble and purified proteins are resolved by SDS-PAGE and detected by Coomassie staining and Western blot hybridization utilizing a Ni-NTA-alkaline phosphatase conjugate (Qiagen) which recognizes his-tagged proteins as described in Example 31(c)(iii). This analysis permits the determination of expression levels of the pHisBotD protein (i.e., number of mg/liter expressed as a soluble protein). The purified BotD protein will migrate as a single band of the predicted MW (i.e., ~50kD).

The level of expression of the pHisBotD protein may be modified (increased) by substitution of the T7 promoter for the T7lac promoter, or by inclusion of the laclq gene on the expression plasmid, and plasmid expressed in BL21(DE3) cell lines in fermentation cultures as described in Example 30. If only very low levels (i.e., less than about 0.5%) of soluble pHisBotD protein are expressed using the above expression systems, the pHisBotD construct may be co-expressed with pACYCGro construct as described in Example 32. In this case, the recombinant BotD protein may co-purify with the folding chaperones. The contaminating chaperones may be removed

as described in Example 34. Preparations of purified pHisBotD protein are tested for endotoxin contamination using the LAL assay as described in Example 24.

The purified pHisBotD protein is used to generate neutralizing antibodies.

BALBc mice are immunized with the BotD protein using Gerbu GMDP adjuvant (CC Biotech) as described in Example 36. The ability of the anti-BotD antibodies to neutralize native C. botulinum type D toxin is demonstrated using the mouse-C. botulinum neutralization model described in Example 36.

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EXAMPLE 48

Expression Of The C Fragment Of The C. botulinum

Serotype F Toxin Gene And Generation Of Neutralizing Antibodies

The C. botulinum type F neurotoxin gene has been cloned and sequenced [East et al. (1992) FEMS Microbiol. Lett. 96:225]. The nucleotide sequence of the toxin gene derived from the 202F strain (ATCC 23387) is available from the EMBL/GenBank sequence data banks under the accession number M92906; the nucleotide sequence of the coding region is listed in SEQ ID NO:70. The amino acid sequence of the C. botulinum type F neurotoxin derived from the 202F strain is listed in SEQ ID NO:71.

The DNA sequence encoding the native *C. botulinum* serotype F C fragment gene derived from the 202F strain can be expressed using the pETHisb vector; the resulting coding region is listed in SEQ ID NO:72 and the corresponding amino acid sequence is listed in SEQ ID NO:73. The C fragment region from any strain of *C. botulinum* serotype F can be amplified and expressed using the approach illustrated below using the C fragment derived from *C. botulinum* type F 202F strain. Expression of the C fragment of *C. botulinum* type F toxin in heterologous hosts (e.g., E. coli) has not been previously reported.

The C fragment of the C. botulinum serotype F (BotF) toxin gene is cloned using the protocols and conditions described in Example 28 for the isolation of the

native BotA gene. The *C. botulinum* type F 202F strain is obtained from the American Type Culture Collection (ATCC 23387). Alternatively, sequences encoding the BotF toxin may be isolated from any BotF expressing strain [e.g., VPI 4404 (ATCC 25764), VPI 2382 (ATCC 27321) and Langeland (ATCC 35415)].

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The following primer pair is used to amplify the BotF gene: 5'-CGCCATGGC TATTCTAATTATATATTTTAATAG-3' [5' primer, engineered NcoI site underlined (SEQ ID NO:74)] and 5'-GCAAGCTTTCATTCTTTCCATCCATTCTC-3' [3' primer, engineered HindIII site underlined, native gene termination codon italicized (SEQ ID NO:75)]. Following PCR amplification, the PCR product is inserted into the pCRscript vector and then the 1.5 kb fragment is cloned into pETHisb vector as described for BotA C fragment gene in Example 28. The resulting construct is termed pHisBotF.

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pHisBotF expresses the BotF gene sequences under the transcriptional control of the T7 lac promoter and the resulting protein contains an N-terminal 10XHis-tag affinity tag. The pHisBotF expression construct is transformed into BL21(DE3) pLysS competent cells and 1 liter cultures are grown, induced and his-tagged proteins are purified utilizing a NiNTA resin as described in Example 28. Total, soluble and purified proteins are resolved by SDS-PAGE and detected by Coomassie staining and Western blot hybridization utilizing a Ni-NTA-alkaline phosphatase conjugate (Qiagen) which recognizes his-tagged proteins as described in Example 31(c)(iii). This analysis permits the determination of expression levels of the pHisBotF protein (i.e., number of mg/liter expressed as a soluble protein). The purified BotF protein will migrate as a single band of the predicted MW (i.e., ~50kD).

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The level of expression of the pHisBotF protein may be modified (increased) by substitution of the T7 promoter for the T7lac promoter, or by inclusion of the lacIq gene on the expression plasmid, and plasmid expressed in BL21(DE3) cell lines in fermentation cultures as described in Example 30. If only very low levels (i.e., less than about 0.5%) of soluble pHisBotF protein are expressed using the above expression systems, the pHisBotF construct may be co-expressed with pACYCGro construct as

described in Example 32. In this case, the recombinant BotF protein may co-purify with the folding chaperones. The contaminating chaperones may be removed as described in Example 34. Preparations of purified pHisBotF protein are tested for endotoxin contamination using the LAL assay as described in Example 24.

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The purified pHisBotF protein is used to generate neutralizing antibodies.

BALBc mice are immunized with the BotF protein using Gerbu GMDP adjuvant (CC Biotech) as described in Example 36. The ability of the anti-BotF antibodies to neutralize native C. botulinum type F toxin is demonstrated using the mouse-C. botulinum neutralization model described in Example 36.

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EXAMPLE 49

Expression Of The C Fragment Of The C. botulinum

Serotype G Toxin Gene And Generation Of Neutralizing Antibodies

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The C. botulinum type G neurotoxin gene has been cloned and sequenced [Campbell et al. (1993) Biochimica et Biophysica Acta 1216:487 and Binz et al. (1990) Nucleic Acids Res. 18:5556]. The nucleotide sequence of the toxin gene derived from the 113/30 strain (NCFB 3012) is available from the EMBL/GenBank sequence data banks under the accession number X74162; the nucleotide sequence of the coding region is listed in SEQ ID NO:76. The amino acid sequence of the C. botulinum type G neurotoxin derived from this strain is listed in SEQ ID NO:77.

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The DNA sequence encoding the native *C. botulinum* serotype G C fragment gene derived from the 113/30 strain can be expressed using the pETHisb vector; the resulting coding region is listed in SEQ ID NO:78 and the corresponding amino acid sequence is listed in SEQ ID NO:79. The C fragment region from any strain of *C. botulinum* serotype G can be amplified and expressed using the approach illustrated below using the C fragment derived from *C. botulinum* type G 113/30 strain.

Expression of the C fragment of *C. botulinum* type G toxin in heterologous hosts (e.g., *E. coli*) has not been previously reported.

The C fragment of the C. botulinum serotype G (BotG) toxin gene is cloned using the protocols and conditions described in Example 28 for the isolation of the native BotA gene. The C. botulinum type G 113/30 strain is obtained from the NCFB. The following primer pair is used to amplify the BotG gene: 5'-CGCCATGGCTGAC ACAATTTTAATACA AGT-3' [5' primer, engineered NcoI site underlined (SEQ ID NO:80)] and 5'-GCCTCGAGTTATTCTGTCCATCCTTCATCCAC-3' [3' primer, engineered XhoI site underlined, native gene termination codon italicized (SEQ ID NO:81)]. Following PCR amplification, the PCR product is inserted into the pCRscript vector and then the 1.5 kb fragment is cloned into pETHisb vector as described for BotA C fragment gene in Example 28 with the exception that the sequences encoding BotG are excised from the pCRscript vector by digestion with NcoI and XhoI and the NcoI site is blunted (the BotG sequences contain an internal HindIII site). This NcoI(filled)/XhoI fragment is then ligated to the pETHisb vector which has been digested with NheI and SalI and the NheI site is blunted. The resulting construct is termed pHisBotG.

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pHisBotG expresses the BotG gene sequences under the transcriptional control of the T7 lac promoter and the resulting protein contains an N-terminal 10XHis-tag affinity tag. The pHisBotG expression construct is transformed into BL21(DE3) pLysS competent cells and 1 liter cultures are grown, induced and his-tagged proteins are purified utilizing a NiNTA resin as described in Example 28. Total, soluble and purified proteins are resolved by SDS-PAGE and detected by Coomassie staining and Western blot hybridization utilizing a Ni-NTA-alkaline phosphatase conjugate (Qiagen) which recognizes his-tagged proteins as described in Example 31(c)(iii). This analysis permits the determination of expression levels of the pHisBotG protein (i.e., number of mg/liter expressed as a soluble protein). The purified BotG protein will migrate as a single band of the predicted MW (i.e., ~50kD).

The level of expression of the pHisBotG protein may be modified (increased) by substitution of the T7 promoter for the T7lac promoter, or by inclusion of the lacIq gene on the expression plasmid, and plasmid expressed in BL21(DE3) cell lines in

fermentation cultures as described in Example 30. If only very low levels (i.e., less than about 0.5%) of soluble pHisBotG protein are expressed using the above expression systems, the pHisBotG construct may be co-expressed with pACYCGro construct as described in Example 32. In this case, the recombinant BotG protein may co-purify with the folding chaperones. The contaminating chaperones may be removed as described in Example 34. Preparations of purified pHisBotG protein are tested for endotoxin contamination using the LAL assay as described in Example 24.

The purified pHisBotG protein is used to generate neutralizing antibodies.

BALBc mice are immunized with the BotG protein using Gerbu GMDP adjuvant (CC Biotech) as described in Example 36. The ability of the anti-BotG antibodies to neutralize native C. botulinum type G toxin is demonstrated using the mouse-C. botulinum neutralization model described in Example 36.

EXAMPLE 50

Expression Of Recombinant Botulinal Toxin Proteins In Eucaryotic Host Cells

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Recombinant botulinal C fragment proteins may be expressed in eucaryotic host cells, such as yeast and insect cells.

a) Expression In Yeast

Botulinal C fragments derived from serotypes A, B, C, D, E, F and G may be expressed in yeast cells using a variety of commercially available vectors. For example, the pPIC3K and pPIC9K expression vectors (Invitrogen) may be employed for expression in the methylotrophic yeast, *Pichia pastoris*. When the pPIC3K vector is employed, expression of the botulinal C fragment protein will be intracellular. When the pPIC3K vector is employed, the botulinal C fragment protein will be secreted (the alpha factor secretion signal is provided on the pPIC9K vector).

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DNA sequences encoding the desired C fragment is inserted into these vectors using techniques known to the art. Briefly, the desired botulinal expression cassette (including sequences encoding the his-tag; described in the preceding examples) is

amplified using the PCR in conjunction with primers that incorporate unique restriction sites at the termini of the amplified fragment. Suitable restriction enzyme sites include SnaBI, EcoRI, AvrII and NotI. When the botulinal C fragment is to be expressed using the pPIC3K vector, the initiator methionine (ATG) is provided by the desired Bot gene sequence and a Kozak consensus sequence is engineered upstream of the ATG (e.g., ACCATGG).

The amplified restriction fragment containing the botulinal C fragment gene is then cloned into the desired expression vector. Recombinant clones are integrated into the *Pichia pastoris* genome and recombinant protein expression is induced using methanol following the manufacturer's instructions (Invitrogen Pichia expression kit manual).

C. botulinum genes are A/T rich and contain multiple sequences that are similar to yeast transcriptional termination signals (e.g., TTTTTATA). If premature transcription termination is observed when the botulinal C fragment genes are expressed in yeast, the transcription termination signals present in the C fragment genes can be removed by either site directed mutagenesis (utilizing the pALTER system; Promega) or by construction of synthetic genes utilizing overlapping synthetic primers.

The botulinal C fragment genes may be expressed in other yeast cells using other commercially available vectors [e.g., using the pYES2 vector (Invitrogen) and S. cerevisiae cells (Invitrogen)].

b) Expression In Insect Cells

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Botulinal C fragments derived from serotypes A, B, C, D, E, F and G may be expressed in insect cells using a variety of commercially available vectors. For example, the pBlueBac4 transfer vector (Invitrogen) may be employed for expression in *Spodoptera frugiperda* (Sf9) insect cells (baculovirus expression system) (equivalent baculovirus vectors and host cells are available from other vendors, e.g., Pharmingen, San Diego, CA). Botulinal C fragments contained on *Ncol/HindIII* fragments

contained within the pHisBotA-G expression constructs (described in the preceding examples) are cloned into the pBlueBac4 vector (digested with Ncol and HindIII); the NcoI site present on the C fragment constructs overlaps with the start codon of the fusion proteins. In the case of botulinal C fragment clones that contain internal HindIII sites (e.g., using the BotG sequences described in Ex. 49), the C fragment gene is contained within a NcoI/XhoI fragment on the pHisBot construct. This Ncol/XhoI fragment is excised from pHisBot and inserted into pBlueBac4 digested with Ncol and Sall. Recombinant baculoviruses are made and the desired recombinant C fragment is expressed in Ss9 cells using the protocols provided by the manufacturer (Invitrogen MaxBac manual). The resulting constructs will express the pHisBot protein intracellularly (including the N-terminal his-tag) under the control of the polyhedrin promoter. For extracellular secretion of botulinal C fragment proteins, the C fragment sequences from the pHisBot constructs are cloned into the pMelBacB vector (Invitrogen) as described above for the pBlueBac4 vector. When the pMelBacB vector is employed, the his-tagged botulinal C fragment proteins are secreted (utilizing a vector-encoded honeybee melittin secretion signal) and contain a nine amino acid extension at the N-terminus.

His-tagged botulinal C fragments expressed in yeast or insect cells are purified using metal chelation columns as described in the preceding examples.

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From the above it is clear that the present invention provides compositions and methods for the preparation of effective multivalent vaccines against *C. botulinum* neurotoxin. It is also contemplated that the recombinant botulinal proteins be used for the production of antitoxins. All publications and patents mentioned in the above specification are herein incorporated by reference. Various modifications and variations of the described method and system of the invention will be apparent to those skilled in the art without departing from the scope and spirit of the invention.

SEQUENCE LISTING

(1) GENERAL INFORMATION:

- (i) APPLICANT: Williams, James A.
 Thalley, Bruce S.
- (ii) TITLE OF INVENTION: Multivalent Vaccine For Clostridium Botulinum Neurotoxin
- (iii) NUMBER OF SEQUENCES: 82
- (iv) CORRESPONDENCE ADDRESS:
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 - (C) CITY: San Francisco
 - (D) STATE: California
 - (E) COUNTRY: United States of America
 - (F) ZIP: 94104
- (v) COMPUTER READABLE FORM:
 - (A) MEDIUM TYPE: Floppy disk
 - (B) COMPUTER: IBM PC compatible
 - (C) OPERATING SYSTEM: PC-DOS/MS-DOS
 - (D) SOFTWARE: PatentIn Release #1.0, Version #1.30
- (vi) CURRENT APPLICATION DATA:
 - (A) APPLICATION NUMBER: US
 - (B) FILING DATE:
 - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
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 - (B) REGISTRATION NUMBER: 40,027
 - (C) REFERENCE/DOCKET NUMBER: OPHD-02304
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 - (B) TELEFAX: (415) 397-8338
- (2) INFORMATION FOR SEQ ID NO:1:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 24 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

GGAAATTTAG CTGCAGCATC TGAC

(2)	INFORMATION FOR SEQ ID NO:2:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 24 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:	
TCT	CAGCAAAT TCGCTTGTGT TGAA	24
(2)	INFORMATION FOR SEQ ID NO:3:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 20 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:	
CTC	GCATATA GCATTAGACC	20
(2)	INFORMATION FOR SEQ ID NO:4:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
,	(ii) MOLECULE TYPE: DNA (genomic)	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:	
CTA	TCTAGGC CTAAAGTAT	19
(2)	INFORMATION FOR SEQ ID NO:5:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 8133 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	(ii) MOLECULE TYPE: DNA (genomic)	
	(ix) FEATURE: (A) NAME/KEY: CDS (B) LOCATION: 18130	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:	
	TCT TTA ATA TCT AAA GAA GAG TTA ATA AAA CTC GCA TAT AGC ATT Ser Leu Ile Ser Lys Glu Glu Leu Ile Lys Leu Ala Tyr Ser Ile 5 10 15	48

	AGA Arg	CCA Pro	AGA Arg	GAA Glu 20	AAT Asn	GAG Glu	TAT Tyr	AAA Lys	ACT Thr 25	ATA Ile	CTA Leu	ACT Thr	AAT Asn	TTA Leu 30	GAC Asp	GAA Glu	96
	TAT Tyr	AAT Asn	AAG Lys 35	TTA Leu	ACT Thr	ACA Thr	AAC Asn	AAT Asn 40	AAT Asn	GAA Glu	AAT Asn	AAA Lys	TAT Tyr 45	TTG Leu	CAA Gln	TTA Leu	144
	AAA Lys	AAA Lys 50	CTA Leu	AAT Asn	GAA Glu	TCA Ser	ATT Ile 55	GAT Asp	GTT Val	TTT Phe	ATG Met	AAT Asn 60	AAA Lys	TAT Tyr	AAA Lys	ACT Thr	192
										CTA Leu							240
										ACA Thr 90							288
										GTC Val							336
•										GCA Ala							384
										AAT Asn							432
										CAG Gln						ATT Ile 160	480
										TTT Phe 170							528
										ATA Ile							576
										GAT Asp							624
										ACT Thr			Glu				672
	ACA Thr 225	AAT Asn	TCT Ser	TTG Leu	AGA Arg	AAA Lys 230	ATA Ile	AAT Asn	AGT Ser	AAT Asn	CAT His 235	GGG Gly	ATA Ile	GAT Asp	ATC Ile	AGG Arg 240	720
	GCT Ala	AAT Asn	AGT Ser	TTG Leu	TTT Phe 245	ACA Thr	GAA Glu	CAA Gln	GAG Glu	TTA Leu 250	TTA Leu	AAT Asn	ATT Ile	TAT Tyr	AGT Ser 255	CAG Gln	768

GAG Glu	TTG Leu	TTA Leu	AAT Asn 260	CGT Arg	GGA Gly	AAT Asn	TTA Leu	GCT Ala 265	GCA Ala	GCA Ala	TCT Ser	GAC Asp	ATA Ile 270	GTA Val	AGA Arg	816
TTA Leu	TTA Leu	GCC Ala 275	CTA Leu	AAA Lys	AAT Asn	TTT Phe	GGC Gly 280	GGA Gly	GTA Val	TAT Tyr	TTA Leu	GAT Asp 285	GTT Val	GAT Asp	ATG Met	864
CTT Leu	CCA Pro 290	GGT Gly	ATT Ile	CAC His	TCT Ser	GAT Asp 295	TTA Leu	TTT Phe	AAA Lys	ACA Thr	ATA Ile 300	TCT Ser	AGA Arg	CCT Pro	AGC Ser	912
TCT Ser 305	ATT Ile	GGA Gly	CTA Leu	GAC Asp	CGT Arg 310	TGG Trp	GAA Glu	ATG Met	ATA Ile	AAA Lys 315	TTA Leu	GAG Glu	GCT Ala	ATT Ile	ATG Met 320	960
AAG Lys	TAT Tyr	AAA Lys	AAA Lys	TAT Tyr 325	ATA Ile	AAT Asn	AAT Asn	TAT Tyr	ACA Thr 330	TCA Ser	GAA Glu	AAC Asn	TTT Phe	GAT Asp 335	AAA Lys	1008
CTT Leu	GAT Asp	CAA Gln	CAA Gln 340	TTA Leu	AAA Lys	GAT Asp	AAT Asn	TTT Phe 345	AAA Lys	CTC. Leu	ATT Ile	ATA Ile	GAA Glu 350	AGT Ser	AAA Lys	1056
AGT Ser	GAA Glu	AAA Lys 355	TCT Ser	GAG Glu	ATA Ile	TTT Phe	TCT Ser 360	AAA Lys	TTA Leu	GAA Glu	AAT Asn	TTA Leu 365	AAT Asn	GTA Val	TCT Ser	1104
Asp	Leu 370	GAA Glu	Ile	Lys	Ile	Ala 375	Phe	Ala	Leu	Gly	Ser 380	Val	Ile	Asn	Gln	1152
Ala 385	Leu	ATA Ile	Ser	Lys	Gln 390	Gly	Ser	Tyr	Leu	Thr 395	Asn	Leu	Val	Ile	Glu 400	1200
Gln	Val	AAA Lys	Asn	Arg 405	Tyr	Gln	Phe	Leu	Asn 410	Gln	His	Leu	Asn	Pro 415	Ala	1248
Ile	Glu	TCT Ser	Asp 420	Asn	Asn	Phe	Thr	Asp 425	Thr	Thr	Lys	Ile	Phe 430	His	Asp	1296
Ser	Leu	TTT Phe 435	Asn	Ser	Ala	Thr	Ala 440	Glu	Asn	Ser	Met	Phe 445	Leu	Thr	Lys	1344
ATA Ile	GCA Ala 450	CCA Pro	TAC Tyr	TTA Leu	CAA Gln	GTA Val 455	GGT Gly	TTT Phe	ATG Met	CCA Pro	GAA Glu 460	GCT Ala	CGC Arg	TCC Ser	ACA Thr	1392
Ile 465	Ser	TTA Leu	Ser	Gly	Pro 470	Gly	Ala	Tyr	Ala	Ser 475	Ala	Ty.r	Tyr	Asp	Phe 480	1440
ATA Ile	AAT Asn	TTA Leu	CAA Gln	GAA Glu 485	Asn	ACT Thr	ATA Ile	GAA Glu	AAA Lys 490	ACT Thr	TTA Leu	AAA Lys	GCA Ala	TCA Ser 495	GAT Asp	1488

		GAA Glu														1536
		ATA Ile 515													TAT Tyr	1584
		GAG Glu														1632
		GGG Gly														1680
		AAT Asn														1728
		TAT Tyr														1776
		GCA Ala 595														1824
		CAA Gln										Tyr				1872
		GGA Gly														1920
		AAA Lys														1968
		GAA Glu														2016
		AAT Asn 675														2064
		AAA Lys														2112
TAT Tyr 705	GAT Asp	TTT Phe	AAT Asn	GTT Val	GAA Glu 710	GAA Glu	ACT Thr	TAT Tyr	CCT Pro	GGG Gly 715	AAG Lys	TTG Leu	CTA Leu	TTA Leu	AGT Ser 720	2160
ATT Ile	ATG Met	GAC Asp	AAA Lys	ATT Ile 725	ACT Thr	TCC Ser	ACT Thr	TTA Leu	CCT Pro 730	GAT Asp	GTA Val	AAT Asn	AAA Lys	AAT Asn 735	TCT Ser	2208

ATT Ile	ACT Thr	ATA Ile	GGA Gly 740	GCA Ala	AAT Asn	CAA Gln	TAT Tyr	GAA Glu 745	GTA Val	AGA Arg	ATT Ile	AAT Asn	AGT Ser 750	GAG Glu	GGA Gly		2256
AGA Arg	AAA Lys	GAA Glu 755	CTT Leu	CTG Leu	GCT Ala	CAC His	TCA Ser 760	GGT Gly	AAA Lys	TGG Trp	ATA Ile	AAT Asn 765	AAA Lys	GAA Glu	GAA Glu		2304
GCT Ala	ATT Ile 770	ATG Met	AGC Ser	GAT Asp	TTA Leu	TCT Ser 775	AGT Ser	AAA Lys	GAA Glu	TAC Tyr	ATT Ile 780	TTT Phe	TTT Phe	GAT Asp	TCT Ser		2352
ATA Ile 785	GAT Asp	AAT Asn	AAG Lys	CTA Leu	AAA Lys 790	GCA Ala	AAG Lys	TCC Ser	AAG Lys	AAT Asn 795	ATT Ile	CCA Pro	GGA Gly	TTA Leu	GCA Ala 800		2400
TCA Ser	ATA Ile	TCA Ser	GAA Glu	GAT Asp 805	ATA Ile	FÅY	ACA Thr	TTA Leu	TTA Leu 810	CTT Leu	GAT Asp	GCA Ala	AGT Ser	GTT Val 815	AGT Ser		2448
CCT Pro	GAT Asp	ACA Thr	AAA Lys 820	TTT Phe	ATT Ile	TTA Leu	AAT Asn	AAT Asn 825	CTT Leu	AAG Lys	CTT Leu	AAT Asn	ATT Ile 830	GAA Glu	TCT Ser		2496
TCT Ser	ATT Ile	GGG Gly 835	GAT Asp	TAC Tyr	ATT Ile	TAT Tyr	TAT Tyr 840	GAA Glu	AAA Lys	TTA Leu	GAG Glu	CCT Pro 845	Val	AAA Lys	AAT Asn		2544
ATA Ile	ATT Ile 850	CAC His	AAT Asn	TCT Ser	ATA Ile	GAT Asp 855	GAT Asp	TTA Leu	ATA Ile	GAT Asp	GAG Glu 860	TTC Phe	AAT Asn	CTA Leu	CTT Leu		2592
GAA Glu 865	AAT Asn	GTA Val	TCT Ser	GAT Asp	GAA Glu 870	TTA Leu	TAT Tyr	GAA Glu	TTA Leu	AAA Lys 875	AAA Lys	TTA Leu	AAT Asn	AAT Asn	CTA Leu 880		2640
GAT Asp	GAG Glu	AAG Lys	TAT Tyr	TTA Leu 885	ATA Ile	TCT Ser	TTT Phe	GAA Glu	GAT Asp 890	ATC Ile	TCA Ser	AAA Lys	AAT Asn	AAT Asn 895	TCA Ser		2688
ACT Thr	TAC Tyr	TCT Ser	GTA Val 900	AGA Arg	TTT Phe	ATT Ile	AAC Asn	AAA Lys 905	AGT Ser	AAT Asn	GGT Gly	GAG Glu	TCA Ser 910	GTT Val	TAT Tyr		2736
GTA Val	GAA Glu	ACA Thr 915	GAA Glu	AAA Lys	GAA Glu	ATT Ile	TTT Phe 920	TCA Ser	AAA Lys	TAT Tyr	AGC Ser	GAA Glu 925	CAT His	ATT Ile	ACA Thr		2784
AAA Lys	GAA Glu 930	ATA Ile	AGT Ser	ACT Thr	ATA Ile	AAG Lys 935	AAT Asn	AGT Ser	ATA Ile	ATT Ile	ACA Thr 940	GAT Asp	GTT Val	AAT Asn	GGT Gly		2832
AAT Asn 945	TTA Leu	TTG Leu	GAT Asp	AAT Asn	ATA Ile 950	CAG Gln	TTA Leu	GAT Asp	CAT His	ACT Thr 955	TCT Ser	CAA Gln	GTT Val	AAT Asn	ACA Thr 960	•	2880
TTA Leu	AAC Asn	GCA Ala	GCA Ala	TTC Phe 965	TTT Phe	ATT Ile	CAA Gln	TCA Ser	TTA Leu 970	ATA Ile	GAT Asp	TAT Tyr	AGT Ser	AGC Ser 975	AAT Asn		2928

AAA GAT GTA CTG AAT GA Lys Asp Val Leu Asn As 980			
GCT CAA CTA TTT AGT AC Ala Gln Leu Phe Ser Th 995			
TTA GTA AAT TTA ATA TO Leu Val Asn Leu Ile Se 1010			
CCT ACA ATA ACA GAG GG Pro Thr Ile Thr Glu Gl 1025			
ATA AAC TTA GGT GCA GC Ile Asn Leu Gly Ala Al 1045		Leu Asp Glu His	
TTA CTA AAA AAA GAA TT Leu Leu Lys Lys Glu Le 1060			Ile Asn
ATG TCA TTA TCT ATA GC Met Ser Leu Ser Ile Al 1075			
GCT GAA GTT ACT ATT TT Ala Glu Val Thr Ile Ph 1090			
ATA CCT TCA TTA GTT AA Ile Pro Ser Leu Val As 1105	n Asn Glu Leu Ile		
TCA GTG GTA AAC TAT TT Ser Val Val Asn Tyr Ph 1125		Glu Ser Lys Lys	
CCT CTT AAA ACA GAA GA Pro Leu Lys Thr Glu As 1140			Asp Leu
GTA ATA TCA GAA ATA GA Val Ile Ser Glu Ile As 1155			
TGT AAT ATA TTA GCA AT Cys Asn Ile Leu Ala Me 1170			
AAT ATA GAT CAC TTT TT Asn Ile Asp His Phe Ph 1185	e Ser Ser Pro Ser	ATA AGT TCT CAT Ile Ser Ser His 1195	ATT CCT 3600 Ile Pro 1200
TCA TTA TCA ATT TAT TC Ser Leu Ser Ile Tyr Se	T GCA ATA GGT ATA	GAA ACA GAA AAT	CTA GAT 3648

TTT Phe	TCA Ser	AAA Lys	AAA Lys 122	Ile	ATG Met	ATG Met	TTA Leu	CCT Pro 122	Asn	GCT Ala	CCT Pro	TCA Ser	AGA Arg 123	Val	TTT Phe	3696
			Thr			GTT Val		Gly					Glu			3744
		Arg				TCA Ser 1255	Ile					Pro				3792
	Trp					TTT Phe					Ile					3840
					Thr	AAT Asn				Lys					Thr	3888
				Met		ACT Thr			Thr					Asn		3936
			Ser			GGA Gly		Gly					Leu			3984
		Tyr				ACG Thr 1335	Asn					Lys				4032
	Ile					AAT Asn					Ile					4080
					Gly	AAG Lys				Asp					Ile	4128
				Asn		CTT Leu			Gly					Asp		4176
TCA Ser	GGC Gly	GAT Asp 1395	Ile	GAT Asp	AAT Asn	AAA Lys	GAT Asp 1400	Arg	TAT Tyr	ATA Ile	TTC Phe	TTG Leu 1405	Thr	TGT Cys	GAG Glu	4224
		Asp				TTA Leu 1415	Ile					Leu				4272
	Tyr					TCT Ser					Tyr					4320
TTA Leu	TCT Ser	AAT Asn	ACT Thr	ATT Ile 1445	Glu	AAA Lys	ATC Ile	AAT Asn	ACT Thr 1450	Leu	GGC Gly	CTA Leu	GAT Asp	AGT Ser 1455	Lys	4368

AAT ATA GCG TAC AAT TAC Asn Ile Ala Tyr Asn Tyr 1460	ACT GAT GAA TCT Thr Asp Glu Ser 1465	AAT AAT AAA TAT Asn Asn Lys Tyr 147	Phe Gly
GCT ATA TCT AAA ACA AGT Ala Ile Ser Lys Thr Ser 1475			
AGT AAA AAT ATA TTA GAA Ser Lys Asn Ile Leu Glu 1490			
AGT AAA GAT TTT ATT GCT Ser Lys Asp Phe Ile Ala 1505	Glu Asp Ile Asn		
ATT AAT ACT ATA ACA GGA Ile Asn Thr Ile Thr Gly 1525		Asp Asn Asn Thr	
AGT ATA GAT TTC TCT ATT Ser Ile Asp Phe Ser Ile 1540			Lys Val
AAT GGA TTA TAT TTA AAT Asn Gly Leu Tyr Leu Asn 1555			
GTG AAA AAT TCA GAT GGA Val Lys Asn Ser Asp Gly 1570			
TTT TTG GAC AAT ATA AGT Phe Leu Asp Asn Ile Ser 1585	Phe Trp Lys Leu		
AAT TTT GTA ATC GAT AAA Asn Phe Val Ile Asp Lys 1605		Val Gly Lys Thr	
GGA TAT GTA GAA TTT ATT Gly Tyr Val Glu Phe Ile 1620			Ile Tyr
TTT GGT GAA TGG AAA ACA Phe Gly Glu Trp Lys Thr 1635	TCG TCA TCT AAA Ser Ser Ser Lys 1640	AGC ACT ATA TTT Ser Thr Ile Phe 1645	AGC GGA 4944 Ser Gly
AAT GGT AGA AAT GTT GTA Asn Gly Arg Asn Val Val 1650			
GAA GAT ATA TCT ACT TCA Glu Asp Ile Ser Thr Ser 1665 1670	Leu Asp Phe Ser	TAT GAA CCT CTC Tyr Glu Pro Leu 1675	TAT GGA 5040 Tyr Gly 1680
ATA GAT AGA TAT ATA AAT Ile Asp Arg Tyr Ile Asn 1685		Ala Pro Asp Leu	

AGT TTA ATA AAT ATT AAT ACC AAT TAT Ser Leu Ile Asn Ile Asn Thr Asn Tyr 1700 170	Tyr Ser Asn Glu Tyr Tyr Pro
GAG ATT ATA GTT CTT AAC CCA AAT ACA	TTC CAC AAA AAA GTA AAT ATA 5184
Glu Ile Ile Val Leu Asn Pro Asn Thr	Phe His Lys Val Asn Ile
1715 1720	1725
AAT TTA GAT AGT TCT TCT TTT GAG TAT	AAA TGG TCT ACA GAA GGA AGT 5232
Asn Leu Asp Ser Ser Ser Phe Glu Tyr	Lys Trp Ser Thr Glu Gly Ser
1730 1735	1740
GAC TTT ATT TTA GTT AGA TAC TTA GAP	GAA AGT AAT AAA AAA ATA TTA 5280
Asp Phe Ile Leu Val Arg Tyr Leu Glu	Glu Ser Asn Lys Lys Ile Leu
1745 1750	1755 1760
CAA AAA ATA AGA ATC AAA GGT ATC TTA	TCT AAT ACT CAA TCA TTT AAT 5328
Gln Lys Ile Arg Ile Lys Gly Ile Leu	Ser Asn Thr Gln Ser Phe Asn
1765	1770 1775
AAA ATG AGT ATA GAT TTT AAA GAT ATT Lys Met Ser Ile Asp Phe Lys Asp Ile 1780 178	Lys Lys Leu Ser Leu Gly Tyr
ATA ATG AGT AAT TTT AAA TCA TTT AAT	TCT GAA AAT GAA TTA GAT AGA 5424
Ile Met Ser Asn Phe Lys Ser Phe Asn	Ser Glu Asn Glu Leu Asp Arg
1795 1800	1805
GAT CAT TTA GGA TTT AAA ATA ATA GAT Asp His Leu Gly Phe Lys Ile Ile Asp 1810 1815	AAT AAA ACT TAT TAC TAT GAT 5472 Asn Lys Thr Tyr Tyr Asp 1820
GAA GAT AGT AAA TTA GTT AAA GGA TTA Glu Asp Ser Lys Leu Val Lys Gly Leu 1825 1830	ATC AAT ATA AAT AAT TCA TTA 5520 Ile Asn Ile Asn Asn Ser Leu 1835 1840
TTC TAT TTT GAT CCT ATA GAA TTT AAC Phe Tyr Phe Asp Pro Ile Glu Phe Asr 1845	TTA GTA ACT GGA TGG CAA ACT Leu Val Thr Gly Trp Gln Thr 1850 1855
ATC AAT GGT AAA AAA TAT TAT TTT GAT Ile Asn Gly Lys Lys Tyr Tyr Phe Asp 1860 186	Ile Asn Thr Gly Ala Ala Leu
ACT AGT TAT AAA ATT ATT AAT GGT AAA	CAC TTT TAT TTT AAT AAT GAT 5664
Thr Ser Tyr Lys Ile Ile Asn Gly Lys	His Phe Tyr Phe Asn Asn Asp
1875 1880	1885
GGT GTG ATG CAG TTG GGA GTA TTT AAA	GGA CCT GAT GGA TTT GAA TAT 5712
Gly Val Met Gln Leu Gly Val Phe Lys	Gly Pro Asp Gly Phe Glu Tyr
1890 1895	1900
TTT GCA CCT GCC AAT ACT CAA AAT AAT	AAC ATA GAA GGT CAG GCT ATA 5760
Phe Ala Pro Ala Asn Thr Gln Asn Asr	Asn Ile Glu Gly Gln Ala Ile
1905 1910	1915 1920
GTT TAT CAA AGT AAA TTC TTA ACT TTC	AAT GGC AAA AAA TAT TAT TTT 5808
Val Tyr Gln Ser Lys Phe Leu Thr Leu	Asn Gly Lys Lys Tyr Tyr Phe
1925	1930 1935

GAT AAT AAC TCA AAA GCA Asp Asn Asn Ser Lys Ala 1940			Asn Glu
AAA TAT TAC TTT AAT CCT Lys Tyr Tyr Phe Asn Pro 1955			
GTA ATT GAC AAT AAT AAG Val Ile Asp Asn Asn Lys 1970			
TCA AAA GGT TGG CAG ACT Ser Lys Gly Trp Gln Thr 1985 1990	Val Asn Gly Ser		
GAT ACC GCT ATT GCC TTT Asp Thr Ala Ile Ala Phe 2005		Thr Ile Asp Gly	
TTT TAT TTT GAT AGT GAT Phe Tyr Phe Asp Ser Asp 2020	TGT GTA GTG AAA Cys Val Val Lys 2025	ATA GGT GTG TTT Ile Gly Val Phe 2030	Ser Thr
TCT AAT GGA TTT GAA TAT Ser Asn Gly Phe Glu Tyr 2035			
ATA GAA GGT CAG GCT ATA Ile Glu Gly Gln Ala Ile 2050	GTT TAT CAA AGT Val Tyr Gln Ser 2055	AAA TTC TTA ACT Lys Phe Leu Thr 2060	TTG AAT 6192 Leu Asn
GGT AAA AAA TAT TAC TTT Gly Lys Lys Tyr Tyr Phe 2065 2070	Asp Asn Asn Ser	AAA GCA GTT ACC Lys Ala Val Thr 2075	GGA TTG 6240 Gly Leu 2080
CAA ACT ATT GAT AGT AAA Gln Thr Ile Asp Ser Lys 2085	AAA TAT TAC TTT Lys Tyr Tyr Phe 209	Asn Thr Asn Thr	GCT GAA 6288 Ala Glu 2095
GCA GCT ACT GGA TGG CAA Ala Ala Thr Gly Trp Gln 2100	ACT ATT GAT GGT Thr Ile Asp Gly 2105	AAA AAA TAT TAC Lys Lys Tyr Tyr 2110	Phe Asn
ACT AAC ACT GCT GAA GCA Thr Asn Thr Ala Glu Ala 2115			
AAA TAT TAC TTT AAT ACT Lys Tyr Tyr Phe Asn Thr 2130	AAC ACT GCT ATA Asn Thr Ala Ile 2135	GCT TCA ACT GGT Ala Ser Thr Gly 2140	TAT ACA 6432 Tyr Thr
ATT ATT AAT GGT AAA CAT Ile Ile Asn Gly Lys His 2145 2150	Phe Tyr Phe Asn	ACT GAT GGT ATT Thr Asp Gly Ile 2155	ATG CAG 6480 Met Gln 2160
ATA GGA GTG TTT AAA GGA Ile Gly Val Phe Lys Gly 2165	CCT AAT GGA TTT Pro Asn Gly Phe 217	Glu Tyr Phe Ala	CCT GCT 6528 Pro Ala 2175

AAT ACG GAT GCT AAC Asn Thr Asp Ala Asn 2180	AAC ATA GAA GGT Asn Ile Glu Gly 218	CAA GCT ATA CTT TAC Gln Ala Ile Leu Tyr 5 219	Gln Asn
GAA TTC TTA ACT TTG Glu Phe Leu Thr Leu 2195			
AAA GCA GTT ACT GGA Lys Ala Val Thr Gly 2210			
AAT CCT AAT AAT GCT Asn Pro Asn Asn Ala 2225	ATT GCT GCA ATT Ile Ala Ala Ile 2230	CAT CTA TGC ACT ATA His Leu Cys Thr Ile 2235	AAT AAT 6720 Asn Asn 2240
GAC AAG TAT TAC TTT Asp Lys Tyr Tyr Phe 2245	Ser Tyr Asp Gly		
ACT ATT GAA AGA AAT . Thr Ile Glu Arg Asn . 2260		Asp Ala Asn Asn Glu	Ser Lys
ATG GTA ACA GGA GTA Met Val Thr Gly Val 2275			
CCT GCT AAT ACT CAC Pro Ala Asn Thr His 2290			
CAG AAC AAA TTC TTA Gln Asn Lys Phe Leu 2305			
GAC TCA AAA GCA GTT ASP Ser Lys Ala Val 2325	Thr Gly Trp Gln		
TAC TTT AAT CTT AAC AT Tyr Phe Asn Leu Asn 2340		Ala Thr Gly Trp Gln	Thr Ile
GAT GGT AAA AAA TAT ' Asp Gly Lys Lys Tyr ' 2355			
GGA TGG CAA ACT ATT (Gly Trp Gln Thr Ile 2 2370			
TTC ATA GCC TCA ACT (Phe Ile Ala Ser Thr (2385			
TTT AAT ACT GAT GGT APPROXIMATION TO THE AST THE ASP Gly 2405	Ile Met Gln Ile		

GGA TTT GAA TAC TTT GCA CCT GCT AAT ACG GAT GCT AAC AAC ATA GAA Gly Phe Glu Tyr Phe Ala Pro Ala Asn Thr Asp Ala Asn Asn Ile Glu 2420 2425 2430	7296
GGT CAA GCT ATA CTT TAC CAA AAT AAA TTC TTA ACT TTG AAT GGT AAA Gly Gln Ala Ile Leu Tyr Gln Asn Lys Phe Leu Thr Leu Asn Gly Lys 2435 2440 2445	7344
AAA TAT TAC TTT GGT AGT GAC TCA AAA GCA GTT ACC GGA CTG CGA ACT Lys Tyr Tyr Phe Gly Ser Asp Ser Lys Ala Val Thr Gly Leu Arg Thr 2450 2455 2460	7392
ATT GAT GGT AAA AAA TAT TAC TTT AAT ACT AAC ACT GCT GTT GCA GTT Ile Asp Gly Lys Lys Tyr Tyr Phe Asn Thr Asn Thr Ala Val Ala Val 2465 2470 2475 2480	7440
ACT GGA TGG CAA ACT ATT AAT GGT AAA AAA TAC TAC TTT AAT ACT AAC Thr Gly Trp Gln Thr Ile Asn Gly Lys Lys Tyr Tyr Phe Asn Thr Asn 2485 2490 2495	7488
ACT TCT ATA GCT TCA ACT GGT TAT ACA ATT ATT AGT GGT AAA CAT TTT Thr Ser Ile Ala Ser Thr Gly Tyr Thr Ile Ile Ser Gly Lys His Phe 2500 2505 2510	7536
TAT TTT AAT ACT GAT GGT ATT ATG CAG ATA GGA GTG TTT AAA GGA CCT Tyr Phe Asn Thr Asp Gly Ile Met Gln Ile Gly Val Phe Lys Gly Pro 2515 2520 2525	7584
GAT GGA TTT GAA TAC TTT GCA CCT GCT AAT ACA GAT GCT AAC AAT ATA Asp Gly Phe Glu Tyr Phe Ala Pro Ala Asn Thr Asp Ala Asn Asn Ile 2530 2540	7632
GAA GGT CAA GCT ATA CGT TAT CAA AAT AGA TTC CTA TAT TTA CAT GAC Glu Gly Gln Ala Ile Arg Tyr Gln Asn Arg Phe Leu Tyr Leu His Asp 2545 2550 2560	7680
AAT ATA TAT TAT TTT GGT AAT AAT TCA AAA GCG GCT ACT GGT TGG GTA Asn Ile Tyr Tyr Phe Gly Asn Asn Ser Lys Ala Ala Thr Gly Trp Val 2565 2570 2575	7728
ACT ATT GAT GGT AAT AGA TAT TAC TTC GAG CCT AAT ACA GCT ATG GGT Thr Ile Asp Gly Asn Arg Tyr Tyr Phe Glu Pro Asn Thr Ala Met Gly 2580 2585 2590	7776
GCG AAT GGT TAT AAA ACT ATT GAT AAT AAA AAT TTT TAC TTT AGA AAT Ala Asn Gly Tyr Lys Thr Ile Asp Asn Lys Asn Phe Tyr Phe Arg Asn 2595 2600 2605	7824
GGT TTA CCT CAG ATA GGA GTG TTT AAA GGG TCT AAT GGA TTT GAA TAC Gly Leu Pro Gln Ile Gly Val Phe Lys Gly Ser Asn Gly Phe Glu Tyr 2610 2615 2620	7872
TTT GCA CCT GCT AAT ACG GAT GCT AAC AAT ATA GAA GGT CAA GCT ATA Phe Ala Pro Ala Asn Thr Asp Ala Asn Asn Ile Glu Gly Gln Ala Ile 2625 2630 2635 2640	7920
CGT TAT CAA AAT AGA TTC CTA CAT TTA CTT GGA AAA ATA TAT TAC TTT Arg Tyr Gln Asn Arg Phe Leu His Leu Leu Gly Lys Ile Tyr Tyr Phe 2645 2650 2655	7968

GGT AAT AAT TCA AAA GCA Gly Asn Asn Ser Lys Ala 2660	GTT ACT GGA TGG C Val Thr Gly Trp G 2665	AA ACT ATT AAT GGT ln Thr Ile Asn Gly 2670	AAA 8016 Lys
GTA TAT TAC TTT ATG CCT Val Tyr Tyr Phe Met Pro 2675	GAT ACT GCT ATG G Asp Thr Ala Met A 2680	CT GCA GCT GGT GGA la Ala Gly Gly 2685	CTT 8064 Leu
TTC GAG ATT GAT GGT GTT Phe Glu Ile Asp Gly Val 2690			
GCC CCT GGG ATA TAT GGC Ala Pro Gly Ile Tyr Gly 2705 271			8133
(2) INFORMATION FOR SEQ	ID NO:6:		
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 2710 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear			
(ii) MOLECULE TYP		2.5	
	CRIPTION: SEQ ID N		-1
Met Ser Leu Ile Ser Lys 1 5	Glu Glu Leu IIe L 10	ys Leu Ala Tyr Ser 15	ile.
Arg Pro Arg Glu Asn Glu 20	Tyr Lys Thr Ile L 25	eu Thr Asn Leu Asp 30	Glu
Tyr Asn Lys Leu Thr Thr 35	Asn Asn Asn Glu A 40	sn Lys Tyr Leu Gln 45	Leu
Lys Lys Leu Asn Glu Ser 50	Ile Asp Val Phe M 55	et Asn Lys Tyr Lys 60	Thr
Ser Ser Arg Asn Arg Ala 65 70		ys Lys Asp Ile Leu 75	Lys 80
Glu Val Ile Leu Ile Lys 85	Asn Ser Asn Thr S	er Pro Val Glu Lys 95	Asn
Leu His Phe Val Trp Ile 100	Gly Gly Glu Val S 105	er Asp Ile Ala Leu 110	Glu
Tyr Ile Lys Gln Trp Ala 115	Asp Ile Asn Ala G 120	lu Tyr Asn Ile Lys 125	Leu
Trp Tyr Asp Ser Glu Ala 130	Phe Leu Val Asn T	hr Leu Lys Lys Ala 140	Ile
Val Glu Ser Ser Thr Thr 145 150			Ile 160
Gln Asn Pro Gln Phe Asp 165	Asn Met Lys Phe T	yr Lys Lys Arg Met 175	Glu
Phe Ile Tyr Asp Arg Gln 180	Lys Arg Phe Ile A	sn Tyr Tyr Lys Ser 190	Gln

Ile Asn Lys Pro Thr Val Pro Thr Ile Asp Asp Ile Ile Lys Ser His Leu Val Ser Glu Tyr Asn Arg Asp Glu Thr Val Leu Glu Ser Tyr Arg Thr Asn Ser Leu Arg Lys Ile Asn Ser Asn His Gly Ile Asp Ile Arg 235 Ala Asn Ser Leu Phe Thr Glu Gln Glu Leu Leu Asn Ile Tyr Ser Gln Glu Leu Leu Asn Arg Gly Asn Leu Ala Ala Ala Ser Asp Ile Val Arg Leu Leu Ala Leu Lys Asn Phe Gly Gly Val Tyr Leu Asp Val Asp Met Leu Pro Gly Ile His Ser Asp Leu Phe Lys Thr Ile Ser Arg Pro Ser Ser Ile Gly Leu Asp Arg Trp Glu Met Ile Lys Leu Glu Ala Ile Met Lys Tyr Lys Lys Tyr Ile Asn Asn Tyr Thr Ser Glu Asn Phe Asp Lys 330 Leu Asp Gln Gln Leu Lys Asp Asn Phe Lys Leu Ile Ile Glu Ser Lys 345 Ser Glu Lys Ser Glu Ile Phe Ser Lys Leu Glu Asn Leu Asn Val Ser 355 Asp Leu Glu Ile Lys Ile Ala Phe Ala Leu Gly Ser Val Ile Asn Gln Ala Leu Ile Ser Lys Gln Gly Ser Tyr Leu Thr Asn Leu Val Ile Glu Gln Val Lys Asn Arg Tyr Gln Phe Leu Asn Gln His Leu Asn Pro Ala Ile Glu Ser Asp Asn Asn Phe Thr Asp Thr Thr Lys Ile Phe His Asp 425 Ser Leu Phe Asn Ser Ala Thr Ala Glu Asn Ser Met Phe Leu Thr Lys Ile Ala Pro Tyr Leu Gln Val Gly Phe Met Pro Glu Ala Arg Ser Thr 455 Ile Ser Leu Ser Gly Pro Gly Ala Tyr Ala Ser Ala Tyr Tyr Asp Phe Ile Asn Leu Gln Glu Asn Thr Ile Glu Lys Thr Leu Lys Ala Ser Asp 490 Leu Ile Glu Phe Lys Phe Pro Glu Asn Asn Leu Ser Gln Leu Thr Glu Gln Glu Ile Asn Ser Leu Trp Ser Phe Asp Gln Ala Ser Ala Lys Tyr 520

- Gln Phe Glu Lys Tyr Val Arg Asp Tyr Thr Gly Gly Ser Leu Ser Glu 530 540

 Asp Asn Gly Val Asp Phe Asn Lys Asn Thr Ala Leu Asp Lys Asn Tyr 545 550 560
- Leu Leu Asn Asn Lys Ile Pro Ser Asn Asn Val Glu Glu Ala Gly Ser 565 570 575
- Lys Asn Tyr Val His Tyr Ile Ile Gln Leu Gln Gly Asp Asp Ile Ser 580 585 590
- Tyr Glu Ala Thr Cys Asn Leu Phe Ser Lys Asn Pro Lys Asn Ser Ile 595 600 605
- Ile Ile Gln Arg Asn Met Asn Glu Ser Ala Lys Ser Tyr Phe Leu Ser 610 615 620
- Asp Asp Gly Glu Ser Ile Leu Glu Leu Asn Lys Tyr Arg Ile Pro Glu 625 630 635 640
- Arg Leu Lys Asn Lys Glu Lys Val Lys Val Thr Phe Ile Gly His Gly
 645 650 655
- Lys Asp Glu Phe Asn Thr Ser Glu Phe Ala Arg Leu Ser Val Asp Ser 660 665 670
- Leu Ser Asn Glu Ile Ser Ser Phe Leu Asp Thr Ile Lys Leu Asp Ile 675 680 685
- Ser Pro Lys Asn Val Glu Val Asn Leu Leu Gly Cys Asn Met Phe Ser 690 695 700
- Tyr Asp Phe Asn Val Glu Glu Thr Tyr Pro Gly Lys Leu Leu Leu Ser 705 710 715 720
- Ile Met Asp Lys Ile Thr Ser Thr Leu Pro Asp Val Asn Lys Asn Ser
- Ile Thr Ile Gly Ala Asn Gln Tyr Glu Val Arg Ile Asn Ser Glu Gly
 740 745 750
- Arg Lys Glu Leu Leu Ala His Ser Gly Lys Trp Ile Asn Lys Glu Glu
 755 760 765
- Ala Ile Met Ser Asp Leu Ser Ser Lys Glu Tyr Ile Phe Phe Asp Ser 770 780
- Ile Asp Asn Lys Leu Lys Ala Lys Ser Lys Asn Ile Pro Gly Leu Ala
 785 790 795 800
- Ser Ile Ser Glu Asp Ile Lys Thr Leu Leu Leu Asp Ala Ser Val Ser 805 810 815
- Pro Asp Thr Lys Phe Ile Leu Asn Asn Leu Lys Leu Asn Ile Glu Ser 820 825 830
- Ser Ile Gly Asp Tyr Ile Tyr Tyr Glu Lys Leu Glu Pro Val Lys Asn 835 840 845
- Ile Ile His Asn Ser Ile Asp Asp Leu Ile Asp Glu Phe Asn Leu Leu 850 855 860

- Glu Asn Val Ser Asp Glu Leu Tyr Glu Leu Lys Lys Leu Asn Asn Leu 865 870 875 880
- Asp Glu Lys Tyr Leu Ile Ser Phe Glu Asp Ile Ser Lys Asn Asn Ser 885 890 895
- Thr Tyr Ser Val Arg Phe Ile Asn Lys Ser Asn Gly Glu Ser Val Tyr
- Val Glu Thr Glu Lys Glu Ile Phe Ser Lys Tyr Ser Glu His Ile Thr 915 920 925
- Lys Glu Ile Ser Thr Ile Lys Asn Ser Ile Ile Thr Asp Val Asn Gly 930 935 940
- Asn Leu Leu Asp Asn Ile Gln Leu Asp His Thr Ser Gln Val Asn Thr 945 950 955 960
- Leu Asn Ala Ala Phe Phe Ile Gln Ser Leu Ile Asp Tyr Ser Ser Asn 965 970 975
- Lys Asp Val Leu Asn Asp Leu Ser Thr Ser Val Lys Val Gln Leu Tyr 980 985 990
- Ala Gln Leu Phe Ser Thr Gly Leu Asn Thr Ile Tyr Asp Ser Ile Gln
 995 1000 1005
- Leu Val Asn Leu Ile Ser Asn Ala Val Asn Asp Thr Ile Asn Val Leu 1010 1015 1020
- Pro Thr Ile Thr Glu Gly Ile Pro Ile Val Ser Thr Ile Leu Asp Gly 1025 1030 1035 1040
- Ile Asn Leu Gly Ala Ala Ile Lys Glu Leu Leu Asp Glu His Asp Pro 1045 1050 1055
- Leu Leu Lys Lys Glu Leu Glu Ala Lys Val Gly Val Leu Ala Ile Asn 1060 1065 1070
- Met Ser Leu Ser Ile Ala Ala Thr Val Ala Ser Ile Val Gly Ile Gly 1075 1080 1085
- Ala Glu Val Thr Ile Phe Leu Leu Pro Ile Ala Gly Ile Ser Ala Gly 1090 1095 1100
- Ile Pro Ser Leu Val Asn Asn Glu Leu Ile Leu His Asp Lys Ala Thr 1105 1110 1115 1120
- Ser Val Val Asn Tyr Phe Asn His Leu Ser Glu Ser Lys Lys Tyr Gly 1125 1130 1135
- Pro Leu Lys Thr Glu Asp Asp Lys Ile Leu Val Pro Ile Asp Asp Leu 1140 1145 1150
- Val Ile Ser Glu Ile Asp Phe Asn Asn Asn Ser Ile Lys Leu Gly Thr 1155 1160 1165
- Cys Asn Ile Leu Ala Met Glu Gly Gly Ser Gly His Thr Val Thr Gly 1170 1175 1180
- Asn Ile Asp His Phe Phe Ser Ser Pro Ser Ile Ser Ser His Ile Pro 1185 1190 1195 1200

- Ser Leu Ser Ile Tyr Ser Ala Ile Gly Ile Glu Thr Glu Asn Leu Asp 1205 1210 1215
- Phe Ser Lys Lys Ile Met Met Leu Pro Asn Ala Pro Ser Arg Val Phe 1220 1225 1230
- Trp Trp Glu Thr Gly Ala Val Pro Gly Leu Arg Ser Leu Glu Asn Asp 1235 1240 1245
- Gly Thr Arg Leu Leu Asp Ser Ile Arg Asp Leu Tyr Pro Gly Lys Phe 1250 1260
- Tyr Trp Arg Phe Tyr Ala Phe Phe Asp Tyr Ala Ile Thr Thr Leu Lys 1265 1270 1275 1280
- Pro Val Tyr Glu Asp Thr Asn Ile Lys Ile Lys Leu Asp Lys Asp Thr 1285 1290 1295
- Arg Asn Phe Ile Met Pro Thr Ile Thr Thr Asn Glu Ile Arg Asn Lys 1300 1305 1310
- Leu Ser Tyr Ser Phe Asp Gly Ala Gly Gly Thr Tyr Ser Leu Leu Leu 1315 1320 1325
- Ser Ser Tyr Pro Ile Ser Thr Asn Ile Asn Leu Ser Lys Asp Asp Leu 1330 1335 1340
- Trp Ile Phe Asn Ile Asp Asn Glu Val Arg Glu Ile Ser Ile Glu Asn 1345 1350 1355 1360
- Gly Thr Ile Lys Lys Gly Lys Leu Ile Lys Asp Val Leu Ser Lys Ile 1365 1370 1375
- Asp Ile Asn Lys Asn Lys Leu Ile Ile Gly Asn Gln Thr Ile Asp Phe 1380 1385 1390
- Ser Gly Asp Ile Asp Asn Lys Asp Arg Tyr Ile Phe Leu Thr Cys Glu 1395 1400 1405
- Leu Asp Asp Lys Ile Ser Leu Ile Ile Glu Ile Asn Leu Val Ala Lys 1410 1415 1420
- Ser Tyr Ser Leu Leu Ser Gly Asp Lys Asn Tyr Leu Ile Ser Asn 1425 1430 1435 1440
- Leu Ser Asn Thr Ile Glu Lys Ile Asn Thr Leu Gly Leu Asp Ser Lys 1445 1450 1455
- Asn Ile Ala Tyr Asn Tyr Thr Asp Glu Ser Asn Asn Lys Tyr Phe Gly
 1460 1465 1470
- Ala Ile Ser Lys Thr Ser Gln Lys Ser Ile Ile His Tyr Lys Lys Asp 1475 1480 1485
- Ser Lys Asn Ile Leu Glu Phe Tyr Asn Asp Ser Thr Leu Glu Phe Asn 1490 1495 1500
- Ser Lys Asp Phe Ile Ala Glu Asp Ile Asn Val Phe Met Lys Asp Asp 1505 1510 1515 1520
- Ile Asn Thr Ile Thr Gly Lys Tyr Tyr Val Asp Asn Asn Thr Asp Lys 1525 1530 1535

- Ser Ile Asp Phe Ser Ile Ser Leu Val Ser Lys Asn Gln Val Lys Val 1540 1550
- Asn Gly Leu Tyr Leu Asn Glu Ser Val Tyr Ser Ser Tyr Leu Asp Phe 1555 1560 1565
- Val Lys Asn Ser Asp Gly His His Asn Thr Ser Asn Phe Met Asn Leu 1570 1575 1580
- Phe Leu Asp Asn Ile Ser Phe Trp Lys Leu Phe Gly Phe Glu Asn Ile 1585 1590 1595 1600
- Asn Phe Val Ile Asp Lys Tyr Phe Thr Leu Val Gly Lys Thr Asn Leu 1605 1610 1615
- Gly Tyr Val Glu Phe Ile Cys Asp Asn Asn Lys Asn Ile Asp Ile Tyr 1620 1625 1630
- Phe Gly Glu Trp Lys Thr Ser Ser Ser Lys Ser Thr Ile Phe Ser Gly
 1635 1640 1645
- Asn Gly Arg Asn Val Val Val Glu Pro Ile Tyr Asn Pro Asp Thr Gly 1650 1660
- Glu Asp Ile Ser Thr Ser Leu Asp Phe Ser Tyr Glu Pro Leu Tyr Gly 1665 1670 1675 1680
- Ile Asp Arg Tyr Ile Asn Lys Val Leu Ile Ala Pro Asp Leu Tyr Thr 1685 1690 1695
- Ser Leu Ile Asn Ile Asn Thr Asn Tyr Tyr Ser Asn Glu Tyr Tyr Pro 1700 1705 1710
- Glu Ile Ile Val Leu Asn Pro Asn Thr Phe His Lys Lys Val Asn Ile 1715 1720 1725
- Asn Leu Asp Ser Ser Ser Phe Glu Tyr Lys Trp Ser Thr Glu Gly Ser 1730 1735 1740
- Asp Phe Ile Leu Val Arg Tyr Leu Glu Glu Ser Asn Lys Lys Ile Leu 1745 · 1750 1755 1760
- Gln Lys Ile Arg Ile Lys Gly Ile Leu Ser Asn Thr Gln Ser Phe Asn 1765 1770 1775
- Lys Met Ser Ile Asp Phe Lys Asp Ile Lys Lys Leu Ser Leu Gly Tyr 1780 1785 1790
- Ile Met Ser Asn Phe Lys Ser Phe Asn Ser Glu Asn Glu Leu Asp Arg 1795 1800 1805
- Asp His Leu Gly Phe Lys Ile Ile Asp Asn Lys Thr Tyr Tyr Asp 1810 1815 1820
- Glu Asp Ser Lys Leu Val Lys Gly Leu Ile Asn Ile Asn Asn Ser Leu 1825 1830 1835 1840
- Phe Tyr Phe Asp Pro Ile Glu Phe Asn Leu Val Thr Gly Trp Gln Thr 1845 1850 1855
- Ile Asn Gly Lys Lys Tyr Tyr Phe Asp Ile Asn Thr Gly Ala Ala Leu 1860 1865 1870

- Thr Ser Tyr Lys Ile Ile Asn Gly Lys His Phe Tyr Phe Asn Asn Asp 1875 1880 1885
- Gly Val Met Gln Leu Gly Val Phe Lys Gly Pro Asp Gly Phe Glu Tyr 1890 1895 1900
- Phe Ala Pro Ala Asn Thr Gln Asn Asn Ile Glu Gly Gln Ala Ile 1905 1910 1915 1920
- Val Tyr Gln Ser Lys Phe Leu Thr Leu Asn Gly Lys Lys Tyr Tyr Phe 1925 1930 1935
- Asp Asn Asn Ser Lys Ala Val Thr Gly Trp Arg Ile Ile Asn Asn Glu 1940 1945 1950
- Lys Tyr Tyr Phe Asn Pro Asn Asn Ala Ile Ala Ala Val Gly Leu Gln
 1955 1960 1965
- Val Ile Asp Asn Asn Lys Tyr Tyr Phe Asn Pro Asp Thr Ala Ile Ile 1970 1975 1980
- Ser Lys Gly Trp Gln Thr Val Asn Gly Ser Arg Tyr Tyr Phe Asp Thr 1985 1990 1995 2000
- Asp Thr Ala Ile Ala Phe Asn Gly Tyr Lys Thr Ile Asp Gly Lys His 2005 2010 2015
- Phe Tyr Phe Asp Ser Asp Cys Val Val Lys Ile Gly Val Phe Ser Thr 2020 2025 2030
- Ser Asn Gly Phe Glu Tyr Phe Ala Pro Ala Asn Thr Tyr Asn Asn Asn 2035 2040 2045
- Ile Glu Gly Gln Ala Ile Val Tyr Gln Ser Lys Phe Leu Thr Leu Asn 2050 2055 2060
- Gly Lys Lys Tyr Tyr Phe Asp Asn Asn Ser Lys Ala Val Thr Gly Leu 2065 2070 2075 2080
- Gln Thr Ile Asp Ser Lys Lys Tyr Tyr Phe Asn Thr Asn Thr Ala Glu 2085 2090 2095
- Ala Ala Thr Gly Trp Gln Thr Ile Asp Gly Lys Lys Tyr Tyr Phe Asn 2100 2105 2110
- Thr Asn Thr Ala Glu Ala Ala Thr Gly Trp Gln Thr Ile Asp Gly Lys 2115 2120 2125
- Lys Tyr Tyr Phe Asn Thr Asn Thr Ala Ile Ala Ser Thr Gly Tyr Thr 2130 2140
- Ile Ile Asn Gly Lys His Phe Tyr Phe Asn Thr Asp Gly Ile Met Gln 2145 2150 2155 2160
- Ile Gly Val Phe Lys Gly Pro Asn Gly Phe Glu Tyr Phe Ala Pro Ala 2165 2170 2175
- Asn Thr Asp Ala Asn Asn Ile Glu Gly Gln Ala Ile Leu Tyr Gln Asn 2180 2185 2190
- Glu Phe Leu Thr Leu Asn Gly Lys Lys Tyr Tyr Phe Gly Ser Asp Ser 2195 2200 2205

- Lys Ala Val Thr Gly Trp Arg Ile Ile Asn Asn Lys Lys Tyr Tyr Phe 2210 2220
- Asn Pro Asn Asn Ala Ile Ala Ala Ile His Leu Cys Thr Ile Asn Asn 2225 2230 2235 2240
- Asp Lys Tyr Tyr Phe Ser Tyr Asp Gly Ile Leu Gln Asn Gly Tyr Ile 2245 2250 2255
- Thr Ile Glu Arg Asn Asn Phe Tyr Phe Asp Ala Asn Asn Glu Ser Lys 2260 2265 2270
- Met Val Thr Gly Val Phe Lys Gly Pro Asn Gly Phe Glu Tyr Phe Ala 2275 2280 2285
- Pro Ala Asn Thr His Asn Asn Asn Ile Glu Gly Gln Ala Ile Val Tyr 2290 2295 2300
- Gln Asn Lys Phe Leu Thr Leu Asn Gly Lys Lys Tyr Tyr Phe Asp Asn 2305 2310 2315 2320
- Asp Ser Lys Ala Val Thr Gly Trp Gln Thr Ile Asp Gly Lys Lys Tyr 2325 2330 2335
- Tyr Phe Asn Leu Asn Thr Ala Glu Ala Ala Thr Gly Trp Gln Thr Ile 2340 2345 2350
- Asp Gly Lys Lys Tyr Tyr Phe Asn Leu Asn Thr Ala Glu Ala Ala Thr 2355 2360 2365
- Gly Trp Gln Thr Ile Asp Gly Lys Lys Tyr Tyr Phe Asn Thr Asn Thr 2370 2380
- Phe Ile Ala Ser Thr Gly Tyr Thr Ser Ile Asn Gly Lys His Phe Tyr 2385 2390 2395 2400
- Phe Asn Thr Asp Gly Ile Met Gln Ile Gly Val Phe Lys Gly Pro Asn 2405 2410 2415
- Gly Phe Glu Tyr Phe Ala Pro Ala Asn Thr Asp Ala Asn Asn Ile Glu 2420 2425 2430
- Gly Gln Ala Ile Leu Tyr Gln Asn Lys Phe Leu Thr Leu Asn Gly Lys 2435 2440 2445
- Lys Tyr Tyr Phe Gly Ser Asp Ser Lys Ala Val Thr Gly Leu Arg Thr 2450 2455 2460
- Ile Asp Gly Lys Lys Tyr Tyr Phe Asn Thr Asn Thr Ala Val Ala Val 2465 2470 2475 2480
- Thr Gly Trp Gln Thr Ile Asn Gly Lys Lys Tyr Tyr Phe Asn Thr Asn 2485 2490 2495
- Thr Ser Ile Ala Ser Thr Gly Tyr Thr Ile Ile Ser Gly Lys His Phe 2500 2505 2510
- Tyr Phe Asn Thr Asp Gly Ile Met Gln Ile Gly Val Phe Lys Gly Pro 2515 2520 2525
- Asp Gly Phe Glu Tyr Phe Ala Pro Ala Asn Thr Asp Ala Asn Asn Ile 2530 2535 2540

Glu Gly Gln Ala Ile Arg Tyr Gln Asn Arg Phe Leu Tyr Leu His Asp 2545 2550 2555

Asn Ile Tyr Tyr Phe Gly Asn Asn Ser Lys Ala Ala Thr Gly Trp Val 2570

Thr Ile Asp Gly Asn Arg Tyr Tyr Phe Glu Pro Asn Thr Ala Met Gly 2585

Ala Asn Gly Tyr Lys Thr Ile Asp Asn Lys Asn Phe Tyr Phe Arg Asn 2600

Gly Leu Pro Gln Ile Gly Val Phe Lys Gly Ser Asn Gly Phe Glu Tyr 2610

Phe Ala Pro Ala Asn Thr Asp Ala Asn Asn Ile Glu Gly Gln Ala Ile 2630 . 2635

Arg Tyr Gln Asn Arg Phe Leu His Leu Leu Gly Lys Ile Tyr Tyr Phe 2645. 2650

Gly Asn Asn Ser Lys Ala Val Thr Gly Trp Gln Thr Ile Asn Gly Lys 2665

Val Tyr Tyr Phe Met Pro Asp Thr Ala Met Ala Ala Ala Gly Gly Leu 2680

Phe Glu Ile Asp Gly Val Ile Tyr Phe Phe Gly Val Asp Gly Val Lys 2695

Ala Pro Gly Ile Tyr Gly . 2710

- (2) INFORMATION FOR SEQ ID NO:7:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 811 amino acids

 - (B) TYPE: amino acid(C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: unknown
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:

Ser Tyr Lys Ile Ile Asn Gly Lys His Phe Tyr Phe Asn Asn Asp Gly

Val Met Gln Leu Gly Val Phe Lys Gly Pro Asp Gly Phe Glu Tyr Phe

Ala Pro Ala Asn Thr Gln Asn Asn Ile Glu Gly Gln Ala Ile Val

Tyr Gln Ser Lys Phe Leu Thr Leu Asn Gly Lys Lys Tyr Tyr Phe Asp 55

Asn Asn Ser Lys Ala Val Thr Gly Trp Arg Ile Ile Asn Asn Glu Lys

Tyr Tyr Phe Asn Pro Asn Asn Ala Ile Ala Ala Val Gly Leu Gln Val 85

Thr Ala Ile Ala Phe Asn Gly Tyr Lys Thr Ile Asp Gly Lys His Phe 130 135 140

Tyr Phe Asp Ser Asp Cys Val Val Lys Ile Gly Val Phe Ser Thr Ser 145 150 155 160

Asn Gly Phe Glu Tyr Phe Ala Pro Ala Asn Thr Tyr Asn Asn Ile 165 170 175

Glu Gly Gln Ala Ile Val Tyr Gln Ser Lys Phe Leu Thr Leu Asn Gly 180 185 190

Lys Lys Tyr Tyr Phe Asp Asn Asn Ser Lys Ala Val Thr Gly Leu Gln
195 200 205

Thr Ile Asp Ser Lys Lys Tyr Tyr Phe Asn Thr Asn Thr Ala Glu Ala 210 215 220

Ala Thr Gly Trp Gln Thr Ile Asp Gly Lys Lys Tyr Tyr Phe Asn Thr 225 230 235 240

Asn Thr Ala Glu Ala Ala Thr Gly Trp Gln Thr Ile Asp Gly Lys Lys 245 250 255

Tyr Tyr Phe Asn Thr Asn Thr Ala Ile Ala Ser Thr Gly Tyr Thr Ile 260 265 270

Ile Asn Gly Lys His Phe Tyr Phe Asn Thr Asp Gly Ile Met Gln Ile 275 280 285

Gly Val Phe Lys Gly Pro Asn Gly Phe Glu Tyr Phe Ala Pro Ala Asn 290 295 300

Thr Asp Ala Asn Asn Ile Glu Gly Gln Ala Ile Leu Tyr Gln Asn Glu 305 310 315 320

Phe Leu Thr Leu Asn Gly Lys Lys Tyr Tyr Phe Gly Ser Asp Ser Lys 325 330 335

Ala Val Thr Gly Trp Arg Ile Ile Asn Asn Lys Lys Tyr Tyr Phe Asn 340 350

Pro Asn Asn Ala Ile Ala Ala Ile His Leu Cys Thr Ile Asn Asn Asp 355 360 365

Lys Tyr Tyr Phe Ser Tyr Asp Gly Ile Leu Gln Asn Gly Tyr Ile Thr 370 375 380

Ile Glu Arg Asn Asn Phe Tyr Phe Asp Ala Asn Asn Glu Ser Lys Met 385 390 395 400

Val Thr Gly Val Phe Lys Gly Pro Asn Gly Phe Glu Tyr Phe Ala Pro 405 410 415

Ala Asn Thr His Asn Asn Asn Ile Glu Gly Gln Ala Ile Val Tyr Gln
420 425 430

Asn Lys Phe Leu Thr Leu Asn Gly Lys Lys Tyr Tyr Phe Asp Asn Asp Ser Lys Ala Val Thr Gly Trp Gln Thr Ile Asp Gly Lys Lys Tyr Tyr Phe Asn Leu Asn Thr Ala Glu Ala Ala Thr Gly Trp Gln Thr Ile Asp Gly Lys Lys Tyr Tyr Phe Asn Leu Asn Thr Ala Glu Ala Ala Thr Gly 490 Trp Gln Thr Ile Asp Gly Lys Lys Tyr Tyr Phe Asn Thr Asn Thr Phe Ile Ala Ser Thr Gly Tyr Thr Ser Ile Asn Gly Lys His Phe Tyr Phe Asn Thr Asp Gly Ile Met Gln Ile Gly Val Phe Lys Gly Pro Asn Gly Phe Glu Tyr Phe Ala Pro Ala Asn Thr Asp Ala Asn Asn Ile Glu Gly 550 Gln Ala Ile Leu Tyr Gln Asn Lys Phe Leu Thr Leu Asn Gly Lys Lys Tyr Tyr Phe Gly Ser Asp Ser Lys Ala Val Thr Gly Leu Arg Thr Ile Asp Gly Lys Lys Tyr Tyr Phe Asn Thr Asn Thr Ala Val Ala Val Thr 600 Gly Trp Gln Thr Ile Asn Gly Lys Lys Tyr Tyr Phe Asn Thr Asn Thr Ser Ile Ala Ser Thr Gly Tyr Thr Ile Ile Ser Gly Lys His Phe Tyr Phe Asn Thr Asp Gly Ile Met Gln Ile Gly Val Phe Lys Gly Pro Asp Gly Phe Glu Tyr Phe Ala Pro Ala Asn Thr Asp Ala Asn Asn Ile Glu Gly Gln Ala Ile Arg Tyr Gln Asn Arg Phe Leu Tyr Leu His Asp Asn Ile Tyr Tyr Phe Gly Asn Asn Ser Lys Ala Ala Thr Gly Trp Val Thr 690 Ile Asp Gly Asn Arg Tyr Tyr Phe Glu Pro Asn Thr Ala Met Gly Ala 715 Asn Gly Tyr Lys Thr Ile Asp Asn Lys Asn Phe Tyr Phe Arg Asn Gly Leu Pro Gln Ile Gly Val Phe Lys Gly Ser Asn Gly Phe Glu Tyr Phe Ala Pro Ala Asn Thr Asp Ala Asn Asn Ile Glu Gly Gln Ala Ile Arg

Tyr Gln Asn Arg Phe Leu His Leu Cly Lys Ile Tyr Tyr Phe Gly
770 780

Asn Asn Ser Lys Ala Val Thr Gly Trp Gln Thr Ile Asn Gly Lys Val 785 790 795 800

Tyr Tyr Phe Met Pro Asp Thr Ala Met Ala Ala 805 810

- (2) INFORMATION FOR SEQ ID NO:8:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 91 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: unknown
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:

Ser Tyr Lys Ile Ile Asn Gly Lys His Phe Tyr Phe Asn Asn Asp Gly
1 5 10 15

Val Met Gln Leu Gly Val Phe Lys Gly Pro Asp Gly Phe Glu Tyr Phe
20 25 30

Ala Pro Ala Asn Thr Gln Asn Asn Ile Glu Gly Gln Ala Ile Val

Tyr Gln Ser Lys Phe Leu Thr Leu Asn Gly Lys Lys Tyr Tyr Phe Asp 50 55 60

Asn Asn Ser Lys Ala Val Thr Gly Trp Arg Ile Ile Asn Asn Glu Lys 70 75 80

Tyr Tyr Phe Asn Pro Asn Asn Ala Ile Ala Ala 85 90

- (2) INFORMATION FOR SEQ ID NO:9:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 7101 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 1..7098
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

ATG AGT TTA GTT AAT AGA AAA CAG TTA GAA AAA ATG GCA AAT GTA AGA
Met Ser Leu Val Asn Arg Lys Gln Leu Glu Lys Met Ala Asn Val Arg

1 10 15

TTT CGT ACT CAA GAA GAT GAA TAT GTT GCA ATA TTG GAT GCT TTA GAA
Phe Arg Thr Gln Glu Asp Glu Tyr Val Ala Ile Leu Asp Ala Leu Glu
20 25 30

48

GA Gl	А ТАЗ 1 Туз	CAT His	s Asr	ATC Met	TC/ Sei	A GAG	AAI Ast	ı Thi	r GTA	A GTG	GAI L Glu	A AAA 1 Lys 45	Ty	TTI Let	A AAA ı Lys	144
TT! Let	A AAA Lys	ASE	T ATA	AAT Asn	' AG1	TTA Leu 55	ACA Thr	GAT	TATT	TAT	TATA	A GAT	י ארצ	TAT	r aaa : Lys	192
AAA Lys 65	ser	GGT	AGA Arg	AAT Asn	AAA Lys	Ala	TTA Leu	AAA Lys	AAA Lys	TTT Phe	Lys	GAA Glu	TAT Tyr	CTA	GTT Val 80	240
ACA Thr	GAA Glu	GTA Val	TTA Leu	GAG Glu 85	CTA Leu	AAG Lys	AAT Asn	AAT Asn	AAT Asn 90	Leu	ACT Thr	CCA Pro	GTT Val	GAG Glu 95	AAA Lys	288
AAT Asn	TTA Leu	CAT His	TTT Phe 100	GTT Val	TGG Trp	ATT Ile	GGA Gly	GGT Gly 105	Gln	ATA Ile	AAT Asn	GAC Asp	ACT Thr 110	Ala	ATT	336
AAT Asn	TAT	ATA Ile 115	Asn	CAA Gln	TGG Trp	AAA Lys	GAT Asp 120	GTA Val	AAT Asn	AGT Ser	GAT Asp	TAT Tyr 125	AAT Asn	GTT Val	AAT Asn	384
GTT Val	TTT Phe 130	Tyr	GAT Asp	AGT Ser	AAT Asn	GCA Ala 135	TTT Phe	TTG Leu	ATA Ile	AAC Asn	ACA Thr 140	TTG Leu	AAA Lys	AAA Lys	ACT Thr	432
GTA Val 145	Val	GAA Glu	TCA Ser	GCA Ala	ATA Ile 150	AAT Asn	GAT Asp	ACA Thr	CTT Leu	GAA Glu 155	TCA Ser	TTT Phe	AGA Arg	GAA Glu	AAC Asn 160	480
TTA Leu	AAT Asn	GAC Asp	CCT Pro	AGA Arg 165	TTT Phe	GAC Asp	TAT Tyr	AAT Asn	AAA Lys 170	TTC Phe	TTC Phe	AGA Arg	AAA Lys	CGT Arg 175	ATG Met	528
GAA Glu	ATA Ile	ATT Ile	TAT Tyr 180	GAT Asp	AAA Lys	CAG Gln	AAA Lys	AAT Asn 185	TTC Phe	ATA Ile	AAC Asn	TAC Tyr	TAT Tyr 190	AAA Lys	GCT Ala	576
CAA Gln	AGA Arg	GAA Glu 195	GAA Glu	AAT Asn	CCT Pro	GAA Glu	CTT Leu 200	ATA Ile	ATT Ile	GAT Asp	GAT Asp	ATT Ile 205	GTA Val	AAG Lys	ACA Thr	624
TAT Tyr	CTT Leu 210	TCA Ser	AAT Asn	GAG Glu	TAT	TCA Ser 215	AAG Lys	GAG Glu	ATA Ile	GAT Asp	GAA Glu 220	CTT Leu	AAT Asn	ACC Thr	TAT Tyr	672
ATT Ile 225	GAA Glu	GAA Glu	TCC Ser	Leu	AAT Asn 230	AAA Lys	ATT Ile	ACA Thr	CAG Gln	AAT Asn 235	AGT Ser	GGA Gly	AAT Asn	GAT Asp	GTT Val 240	720
AGA Arg	AAC Asn	TTT Phe	Glu	GAA Glu 245	TTT Phe	AAA . Lys .	AAT Asn	GGA Gly	GAG Glu 250	TCA Ser	TTC Phe	AAC Asn	TTA Leu	TAT Tyr 255	GAA Glu	768
CAA Gln	GAG Glu	Leu	GTA Val 260	GAA . Glu .	AGG Arg	TGG .	Asn	TTA Leu 265	GCT Ala	GCT Ala	GCT Ala	TCT (GAC Asp 270	ATA Ile	TTA Leu	816
AGA Arg	ATA Ile	TCT Ser	GCA '	TTA . Leu :	AAA Lys	GAA Glu	ATT Ile	GGT Gly	GGT Gly	ATG Met	TAT Tyr	TTA (GAT Asp	GTT Val	GAT Asp	864

275 280 285

ATG Met	TTA Leu 290	Pro	GGA Gly	ATA Ile	CAA Gln	CCA Pro 295	Asp	TTA Leu	TTT Phe	GAG Glu	TCT Ser 300	Ile	GAG Glu	AAA Lys	CCT	912
AGT Ser 305	Ser	GTA Val	ACA Thr	GTG Val	GAT Asp 310	TTT Phe	TGG Trp	GAA Glu	ATG Met	ACA Thr 315	AAG Lys	TTA Leu	GAA Glu	GCT Ala	ATA Ile 320	960
		TAC Tyr														1008
ATG Met	TTA Leu	GAC Asp	GAA Glu 340	GAA Glu	GTT Val	CAA Gln	AGT Ser	AGT Ser 345	Phe	GAA Glu	TCT Ser	GTT Val	CTA Leu 350	GCT Ala	TCT Ser	1056
AAG Lys	TCA Ser	GAT Asp 355	AAA Lys	TCA Ser	GAA Glu	ATA Ile	TTC Phe 360	TCA Ser	TCA Ser	CTT Leu	GGT Gly	GAT Asp 365	ATG Met	GAG Glu	GCA Ala	1104
		CTA Leu														1152
		CTA Leu		Ser												1200
AAA Lys	CAA Gln	ATC Ile	GAG Glu	AAT Asn 405	AGA Arg	TAT Tyr	AAA Lys	ATA Ile	TTG Leu 410	AAT Asn	AAT Asn	AGT Ser	TTA Leu	AAT Asn 415	CCA Pro	1248
		AGC Ser														1296
		ATA Ile 435														1344
Glu		GGA Gly														1392
ACT Thr 465	ATT Ile	AAC Asn	TTA Leu	AGT Ser	GGC Gly 470	CCT Pro	GAA Glu	GCA Ala	TAT Tyr	GCG Ala 475	GCA Ala	GCT Ala	TAT Tyr	CAA Gln	GAT Asp 480	1440
		ATG Met														1488
		AGA Arg														1536
		GAA Glu 515														1584

GC! Ala	CAL Gli 530	1 Pile	GAA Glu	GAA Glu	TAI	Lys 535	Arg	AA? Asi	r TAT	r TTT	GAA Glu 540	ı Gl	r TC1	CTT Let	GGT Gly	1632
GA/ Glu 545	r war	GAT Asp	AAT Asn	CTI Leu	GAT Asp 550	Phe	TCT Ser	CAF Glr	AAT Asn	TATA	: Val	A GTT	GAC Asp	AAC Lys	GAG Glu 560	1680
TAT Tyr	CTI Lev	TTA Leu	GAA Glu	Lys 565	He	TCT Ser	TCA Ser	TTA Leu	GCA Ala 570	Arg	AGT Ser	TCA Ser	GAG Glu	AGA Arg 575	GGA	1728
TAT	Ile	CAC His	TAT Tyr 580	ATT Ile	GTT Val	CAG Gln	TTA Leu	CAA Gln 585	Gly	GAT Asp	AAA Lys	ATI	Ser 590	TAT Tyr	GAA Glu	1776
GCA Ala	GCA Ala	TGT Cys 595	Asn	TTA Leu	TTT Phe	GCA Ala	AAG Lys 600	ACT Thr	CCT Pro	TAT Tyr	GAT Asp	AGT Ser 605	Val	CTG Leu	Phe	1824
CAG Gln	AAA Lys 610	AAT Asn	ATA Ile	GAA Glu	GAT Asp	TCA Ser 615	GAA Glu	ATT	GCA Ala	TAT Tyr	TAT Tyr 620	TAT Tyr	AAT	CCT Pro	GGA Gly	1872
GAT Asp 625	GTA	GAA Glu	ATA Ile	CAA Gln	GAA Glu 630	ATA Ile	GAC Asp	AAG Lys	TAT Tyr	AAA Lys 635	ATT Ile	CCA Pro	AGT Ser	ATA Ile	ATT Ile 640	1920
TCT Ser	GAT Asp	AGA Arg	CCT Pro	AAG Lys 645	ATT Ile	AAA Lys	TTA Leu	ACA Thr	TTT Phe 650	ATT Ile	GGT Gly	CAT His	GGT Gly	AAA Lys 655	GAT Asp	1968
GAA Glu	TTT Phe	AAT Asn	ACT Thr 660	GAT Asp	ATA Ile	TTT Phe	GCA Ala	GGT Gly 665	TTT Phe	GAT Asp	GTA Val	GAT Asp	TCA Ser 670	TTA Leu	TCC Ser	2016
ACA Thr	GAA Glu	ATA Ile 675	GAA Glu	GCA Ala	GCA Ala	ATA Ile	GAT Asp 680	TTA Leu	GCT Ala	AAA Lys	GAG Glu	GAT Asp 685	ATT Ile	TCT Ser	CCT Pro	2064
AAG Lys	TCA Ser 690	ATA Ile	GAA Glu	ATA Ile	AAT Asn	TTA Leu 695	TTA Leu	GGA Gly	TGT Cys	AAT Asn	ATG Met 700	TTT Phe	AGC Ser	TAC Tyr	TCT Ser	2112
ATC Ile 705	ASI	GTA Val	GAG Glu	GAG Glu	ACT Thr 710	TAT Tyr	CCT Pro	GGA Gly	AAA Lys	TTA Leu 715	Leu	CTT Leu	AAA Lys	GTT Val	AAA Lys 720	2160
GAT Asp	AAA Lys	ATA Ile	Ser	GAA Glu 725	TTA Leu	ATG Met	CCA Pro	TCT Ser	ATA Ile 730	AGT Ser	CAA Gln	GAC Asp	Ser	ATT Ile 735	ATA Ile	2208
GTA Val	AGT Ser	GCA Ala	AAT Asn 740	CAA Gln	TAT Tyr	GAA (Glu	Val .	AGA Arg 745	ATA Ile	AAT Asn	AGT Ser	GAA Glu	GGA Gly 750	AGA Arg	AGA Arg	2256
GAA Glu	TTA Leu	TTG Leu 755	GAT Asp	CAT His	TCT Ser	Gly (GAA ' Glu ' 760	TGG Trp	ATA . Ile .	AAT . Asn	Lys	GAA Glu 765	ĠAA . Glu	AGT Ser	ATT Ile	2304

ATA Ile	AAG Lys 770	Asp	ATT	TCA Ser	TCA Ser	AAA Lys 775	GAA Glu	TAT	ATA Ile	TCA Ser	Phe	Asn	CCT Pro	AAA Lys	GAA Glu	2352
AAT Asn 785	Lys	ATT	ACA Thr	GTA Val	AAA Lys 790	TCT Ser	AAA Lys	AAT Asn	TTA Leu	CCT Pro 795	Glu	CTA Leu	TCT Ser	ACA Thr	TTA Leu 800	2400
TTA Leu	CAA Gln	GAA Glu	ATT	AGA Arg 805	Asn	AAT Asn	TCT Ser	AAT Asn	TCA Ser 810	AGT Ser	GAT Asp	ATT Ile	GAA Glu	CTA Leu 815	GAA Glu	2448
GAA Glu	AAA Lys	GTA Val	ATG Met 820	TTA Leu	ACA Thr	GAA Glu	TGT Cys	GAG Glu 825	ATA Ile	AAT Asn	GTT Val	ATT Ile	TCA Ser 830	AAT Asn	ATA Ile	2496
GAT Asp	ACG Thr	CAA Gln 835	ATT	GTT Val	GAG Glu	GAA Glu	AGG Arg 840	ATT	GAA Glu	GAA Glu	GCT Ala	AAG Lys 845	AAT Asn	TTA Leu	ACT Thr	2544
TCT Ser	GAC Asp 850	TCT Ser	ATT Ile	AAT Asn	TAT Tyr	ATA Ile 855	AAA Lys	GAT Asp	GAA Glu	TTT Phe	AAA Lys 860	CTA Leu	ATA Ile	GAA Glu	TCT Ser	2592
ATT Ile 865	TCT Ser	GAT Asp	GCA Ala	CTA Leu	TGT Cys 870	GAC Asp	TTA Leu	AAA Lys	CAA Gln	CAG Gln 875	AAT Asn	GAA Glu	TTA Leu	GAA Glu	GAT Asp 880	2640
TCT Ser	CAT His	TTT Phe	ATA Ile	TCT Ser 885	TTT Phe	GAG Glu	GAC Asp	ATA Ile	TCA Ser 890	GAG Glu	ACT Thr	GAT Asp	GAG Glu	GGA Gly 895	TTT Phe	2688
												ATA Ile				2736
Thr	Glu	Lys 915	Thr	Ile	Phe	Ser	Glu 920	Tyr	Ala	Asn	His	ATA Ile 925	Thr	Glu	Glu	2784
ATT Ile	TCT Ser 930	AAG Lys	ATA Ile	AAA Lys	GGT Gly	ACT Thr 935	ATA Ile	TTT	GAT Asp	ACT Thr	GTA Val 940	AAT Asn	GGT Gly	AAG Lys	TTA Leu	2832
GTA Val 945	Lys	AAA Lys	GTA Val	AAT Asn	TTA Leu 950	GAT Asp	ACT Thr	ACA Thr	CAC His	GAA Glu 955	GTA Val	AAT Asn	ACT Thr	TTA Leu	AAT Asn 960	2880
GCT Ala	GCA Ala	TTT Phe	TTT Phe	ATA Ile 965	CAA Gln	TCA Ser	TTA Leu	ATA Ile	GAA Glu 970	TAT Tyr	AAT Asn	AGT Ser	TCT Ser	AAA Lys 975	GAA Glu	2928
TCT Ser	CTT Leu	AGT Ser	AAT Asn 980	TTA Leu	AGT Ser	GTA Val	GCÀ Ala	ATG Met 985	AAA Lys	GTC Val	CAA Gln	GTT Val	TAC Tyr 990	GCT Ala	CAA Gln	2976
						Asn		Ile			Ala	GCC Ala 1005	Lys			3024

GAA TTA GTA TCA ACT GCA	TTA GAT GAA ACT ATA	GAC TTA CTT CCT ACA 3072
Glu Leu Val Ser Thr Ala	Leu Asp Glu Thr Ile	Asp Leu Leu Pro Thr
1010	1015	1020
TTA TCT GAA GGA TTA CCT Leu Ser Glu Gly Leu Pro 1025	Ile Ile Ala Thr Ile	Ile Asp Gly Val Ser
TTA GGT GCA GCA ATC AAA	GAG CTA AGT GAA ACG	AGT GAC CCA TTA TTA 3168
Leu Gly Ala Ala Ile Lys	Glu Leu Ser Glu Thr	Ser Asp Pro Leu Leu
1045	1050	1055
AGA CAA GAA ATA GAA GCT	AAG ATA GGT ATA ATG	GCA GTA AAT TTA ACA 3216
Arg Gln Glu Ile Glu Ala	Lys Ile Gly Ile Met	Ala Val Asn Leu Thr
1060	1065	1070
ACA GCT ACA ACT GCA ATC Thr Ala Thr Thr Ala Ile 1075	ATT ACT TCA TCT TTG Ile Thr Ser Ser Leu 1080	GGG ATA GCT AGT GGA 3264 Gly Ile Ala Ser Gly 1085
TTT AGT ATA CTT TTA GTT	CCT TTA GCA GGA ATT	TCA GCA GGT ATA CCA 3312
Phe Ser Ile Leu Leu Val	Pro Leu Ala Gly Ile	Ser Ala Gly Ile Pro
1090	1095	1100
AGC TTA GTA AAC AAT GAA Ser Leu Val Asn Asn Glu 1105 1110	Leu Val Leu Arg Asp	Lys Ala Thr Lys Val
GTA GAT TAT TTT AAA CAT	GTT TCA TTA GTT GAA	ACT GAA GGA GTA TTT 3408
Val Asp Tyr Phe Lys His	Val Ser Leu Val Glu	Thr Glu Gly Val Phe
1125	1130	1135
ACT TTA TTA GAT GAT AAA	ATA ATG ATG CCA CAA	GAT GAT TTA GTG ATA 3456
Thr Leu Leu Asp Asp Lys	Ile Met Met Pro Gln	Asp Asp Leu Val Ile
1140	1145	1150
TCA GAA ATA GAT TTT AAT .	AAT AAT TCA ATA GTT	TTA GGT AAA TGT GAA 3504
Ser Glu Ile Asp Phe Asn .	Asn Asn Ser Ile Val	Leu Gly Lys Cys Glu
1155	1160	1165
ATC TGG AGA ATG GAA GGT of the Trp Arg Met Glu Gly of 1170	GGT TCA GGT CAT ACT Gly Ser Gly His Thr 1175	GTA ACT GAT GAT ATA 3552 Val Thr Asp Asp Ile 1180
GAT CAC TTC TTT TCA GCA GASP His Phe Phe Ser Ala 1185	CCA TCA ATA ACA TAT Pro Ser Ile Thr Tyr 1195	Arg Glu Pro His Leu
TCT ATA TAT GAC GTA TTG (Ser Ile Tyr Asp Val Leu (1205	GAA GTA CAA AAA GAA Glu Val Gln Lys Glu 1210	GAA CTT GAT TTG TCA 3648 Glu Leu Asp Leu Ser 1215
AAA GAT TTA ATG GTA TTA C	CCT AAT GCT CCA AAT	AGA GTA TTT GCT TGG 3696
Lys Asp Leu Met Val Leu 1	Pro Asn Ala Pro Asn	Arg Val Phe Ala Trp
1220	1225	1230
GAA ACA GGA TGG ACA CCA (Glu Thr Gly Trp Thr Pro (1235		

AA? Lys	Lei 125	ı rec	GAC Asp	CGT	ATA	AGA Arg 125	Asp	AAC Asr	TAT Tyz	GAA Glu	GG7 Gly 126	/ Glu	TTI Phe	TAT	TGG		3792
AGA Arg 126	TY	TTT Phe	GCT Ala	TTT	ATA Ile 127	Ala	GAT Asp	GCT Ala	TTA Leu	ATA Ile 127	Thr	A ACA	TTA Leu	AAA Lys	CCA Pro 1280		3840
AGA Arg	TAT	GAA Glu	GAT Asp	ACT Thr 128	Asn	ATA Ile	AGA Arg	ATA	AAT Asn 129	Leu	GAT Asp	AGT Ser	AAT Asn	ACT Thr 129	AGA Arg 5		3888
AGT Ser	TTI Phe	ATA : Ile	GTT Val 130	Pro	ATA Ile	ATA Ile	ACT Thr	ACA Thr 130	Glu	TAT Tyr	ATA Ile	AGA Arg	GAA Glu 131	Lys	TTA		3936
TCA Ser	TAT Tyr	Ser	Phe	TAT	GGT Gly	TCA Ser	GGA Gly 132	Gly	ACT Thr	TAT Tyr	GCA Ala	TTG Leu 132	Ser	CTT Leu	TCT Ser		3984
CAA Gln	TAT Tyr 133	AAT Asn 0	ATG Met	GGT Gly	ATA Ile	AAT Asn 1339	Ile	GAA Glu	TTA Leu	AGT Ser	GAA Glu 134	Ser	GAT Asp	GTT Val	TGG Trp		4032
ATT Ile 134	Ile	GAT Asp	GTT Val	GAT Asp	AAT Asn 1350	Val	GTG Val	AGA Arg	GAT Asp	GTA Val 1355	Thr	ATA Ile	GAA Glu	TCT Ser	GAT Asp 1360		4080
AAA Lys	ATT Ile	AAA Lys	AAA Lys	GGT Gly 1365	Asp	TTA Leu	ATA Ile	GAA Glu	GGT Gly 1370	Ile	TTA Leu	TCT	ACA Thr	CTA Leu 1379	Ser		4128
ATT Ile	GAA Glu	GAG Glu	AAT Asn 1380	Lys	ATT Ile	ATC Ile	TTA Leu	AAT Asn 1389	Ser	CAT His	GAG Glu	ATT Ile	AAT Asn 1390	Phe	TCT Ser		4176
GGT Gly	GAG Glu	GTA Val 1399	Asn	GGA Gly	AGT Ser	AAT Asn	GGA Gly 1400	Phe	GTT Val	TCT	TTA Leu	ACA Thr 1405	Phe	TCA Ser	ATT Ile		4224
TTA Leu	GAA Glu 1410	GGA Gly)	ATA Ile	AAT Asn	Ala	ATT Ile 1415	Ile	GAA Glu	GTT Val	GAT Asp	TTA Leu 1420	Leu	TCT Ser	AAA Lys	TCA Ser		4272
TAT Tyr 1425	Lys	TTA Leu	CTT Leu	Ile	TCT Ser 1430	Gly	GAA Glu	TTA Leu	AAA Lys	ATA Ile 1435	Leu	ATG Met	TTA Leu	AAT Asn	TCA Ser 1440		4320
AAT Asn	CAT His	ATT Ile	Gln	CAG Gln 1445	AAA Lys	ATA Ile	GAT Asp	TAT Tyr	ATA Ile 1450	Gly	TTC Phe	AAT Asn	Ser	GAA Glu 1455	Leu	•	4368
CAG Gln	AAA Lys	AAT Asn	ATA Ile 1460	CCA ' Pro '	TAT Tyr	AGC Ser	Phe	GTA Val 1465	Asp	AGT Ser	GAA Glu	GGA Gly	AAA Lys 1470	Glu	AAT Asn	4	4416
GGT Gly	TTT Phe	ATT Ile 1475	Asn	GGT '	TCA . Ser '	Thr	AAA Lys (1480	GAA Glu	GGT Gly	TTA Leu	TTT Phe	GTA Val 1485	Ser	GAA Glu	TTA Leu	4	1464

CC'	T GA O As 14	p va.	A GTT l Val	CTI Let	T ATA	AGT Ser 149	Lys	GTT Val	TAT Tyr	ATC Met	GA' As ₁	p Asp	AGI Ser	'AAG 'Lys	CCT Pro		4512
TCI Sei 150	c Phe	r GG/ e Gly	A TAI	TAT	C AGT Ser 151	Asn	AAT Asn	TTG Leu	Lys	GAT Asp 151	Va]	C AAA L Lys	GTT Val	ATA Ile	ACT Thr 1520		4560
AA/ Lys	A GAT	TAAT PAST	GTT Val	AAT Asn 152	Ile	TTA Leu	ACA Thr	GGT	TAT Tyr 153	Tyr	CT7	T AAG 1 Lys	GAT Asp	GAT Asp 153			4608
ьys	; 11 6	e Ser	154	Ser 0	Leu	Thr	Leu	Gln 154	Asp 5	Glu	Lys	Thr	11e	Lys 0			4656
ASI	. ser	155	. H1S	Leu	qzA	GLu	156	Gly 0	Val	Ala	Glu	11e	Leu 5	Lys			4704
ATG Met	AAT Asn 157	Arg	Lys	GGT Gly	AAT Asn	ACA Thr 1579	Asn	ACT Thr	TCA Ser	GAT Asp	TCT Ser 158	TTA Leu 0	ATG Met	AGC Ser	TTT Phe		4752
TTA Leu 158	Glu	AGT Ser	ATG Met	AAT Asn	ATA Ile 1590	Lys	AGT Ser	ATT Ile	TTC Phe	GTT Val 1595	Asn	TTC Phe	TTA Leu	CAA Gln	TCT Ser 1600		4800
Asn	IIe	Lys	Phe	Ile 1609	Leu.	Asp	Ala	Asn	Phe 1610	Ile	Ile	AGT Ser	Gly	Thr 1619	Thr	•	4848
ser	TIE	GIA	1620	Phe	Glu	Phe	Ile	Cys 1625	Asp	Glu	Asn	GAT Asp	Asn 1630	Ile	Gln	4	1896
Pro	Tyr	Phe 163	Ile	Lys	Phe	Asn	Thr 1640	Leu	Glu	Thr	Asn	TAT Tyr 1645	Thr	Leu	Tyr	4	1944
Val	Gly 1650	Asn O	Arg	Gln	Asn	Met 1655	Ile	Val	Glu	Pro	Asn 1660		Asp	Leu	Asp	4	1992
1665	ser	GIY	Asp	IIe	Ser 1670	Ser	Thr	Val	Ile	Asn 1675	Phe	TCT Ser	Gln	Lys	Tyr 1680	5	040
CTT Leu	TAT Tyr	GGA Gly	Ile	GAC Asp 1685	Ser	TGT Cys	GTT .	Asn	AAA Lys 1690	Val	GTA Val	ATT Ile	Ser	CCA Pro 1695	Asn	5	880
Ile	Tyr	Thr	Asp 1700	Glu	Ile	Asn	Ile	Thr 1705	Pro	Val '	Tyr		Thr . 1710	Asn	Asn	5	136
ACT Thr	TAT Tyr	CCA Pro 1715	Glu	GTT . Val	ATT Ile	Val :	TTA (Leu) 1720	GAT (Asp	GCA . Ala .	AAT ' Asn '	TAT Tyr	ATA Ile 1725	AAT (Asn (GAA Glu	AAA Lys	5	184

AT	A AA' e Ası 17	ı vaı	AAT Asn	ATC Ile	AAT Asn	GAT Asp 173	Leu	TCT Ser	TATA	A CGA e Arg	TA'	r Val	TGG Tr	AG1 Ser	AAT Asn	52	232
GA? Asp 174	י פדי	r aat / Asn	GAT Asp	TTT	ATT Ile 175	Leu	ATG Met	TCA Ser	ACT Thr	AGT Ser 175	Gl	A GAA 1 Glu	AA1 Asn	'AAG	GTG Val 1760	52	280
361	. GII	A GTT 1 Val	гÀг	11e 176	Arg 5	Phe	Val	Asn	Val 177	Phe 0	Lys	asp	Lys	Thr 177	Leu 5	53	128
GCA Ala	LAA 1 Laan	'AAG Lys	CTA Leu 178	Ser	TTT Phe	AAC Asn	TTT Phe	AGT Ser 178	Asp	AAA Lys	CA/ Glr	A GAT Asp	GTA Val 179	Pro	GTA Val	53	76
AGT Ser	GAA Glu	ATA Ile 179	TTE	TTA Leu	TCA Ser	TTT Phe	ACA Thr 180	Pro	TCA Ser	TAT Tyr	TAT	GAG Glu 180	Asp	GGA Gly	TTG Leu	54	24
ATT Ile	GGC Gly 181	TAT Tyr 0	GAT Asp	TTG Leu	GGT Gly	CTA Leu 1819	Val	TCT	TTA Leu	TAT Tyr	AAT Asn 182	Glu	AAA Lys	TTT Phe	TAT Tyr	54	72
ATT Ile 182	Asn	AAC Asn	TTT Phe	GGA Gly	ATG Met 1830	Met	GTA Val	TCT Ser	GGA Gly	TTA Leu 1835	Ile	TAT Tyr	ATT Ile	AAT Asn	GAT Asp 1840	55:	20
TCA Ser	TTA Leu	TAT	Tyr	TTT Phe 1849	Lys	CCA Pro	CCA Pro	GTA Val	AAT Asn 1850	Asn	TTG Leu	ATA Ile	ACT Thr	GGA Gly 1859	Phe	556	68
GTG Val	ACT	GTA Val	GGC Gly 1860	Asp	GAT Asp	AAA Lys	TAC Tyr	TAC Tyr 1865	Phe	AAT Asn	CCA Pro	ATT Ile	AAT Asn 1870	Gly	GGA Gly	561	16
GCT Ala	GCT Ala	TCA Ser 1875	Ile	GGA Gly	GAG Glu	ACA Thr	ATA Ile 1880	Ile	GAT Asp	GAC Asp	AAA Lys	AAT Asn 1885	Tyr	TAT Tyr	TTC Phe	566	54
Asn	1890		Gly	Val	Leu	Gln 1895	Thr	Gly	Val	Phe	Ser 1900	Thr	Glu	Asp	Gly	571	L2
TTT Phe 1909	Lys	TAT Tyr	TTT (Ala	CCA Pro 1910	Ala	AAT Asn	ACA Thr	Leu	GAT Asp 1915	Glu	AAC Asn	CTA Leu	Glu	GGA Gly 1920	576	0
GAA Glu	GCA Ala	ATT Ile	Asp :	TTT Phe 1925	Thr	GGA . Gly	AAA Lys	Leu	ATT Ile 1930	Ile	GAC Asp	GAA Glu	Asn	ATT Ile 1935	Tyr	580	8
TAT Tyr	TTT Phe	GAT Asp	GAT A Asp A	AAT Asn	TAT .	AGA (Arg (Gly A	GCT Ala 1945	GTA Val	GAA Glu	TGG Trp	Lys	GAA Glu 1950	TTA Leu	GAT Asp	585	6
GGT Gly	GAA Glu	ATG Met 1955	CAC 1	rat ' Cyr	TTT I	Ser	CCA (Pro (1960	GAA / Glu '	ACA Thr	GGT . Gly :	AAA Lys	GCT Ala 1965	TTT . Phe	AAA Lys	GGT Gly	590	4

CT <i>I</i> Let	A AAT 1 Asi 197	1 GII	A ATA	GGT Gly	GAT Asp	TAT Tyr 197	Lys	TAC Tyr	TAT	TTC Phe	AA? Asr 198	ı Ser	GAT Asp	GGA Gly	GTT Val	5952
ATO Met 198	GII	A AAA 1 Lys	GGA Gly	TTT	GTT Val 199	Ser	ATA Ile	AAT Asn	GAT Asp	AAT Asn 199	Lys	CAC His	TAT Tyr	TTT Phe	GAT Asp 2000	6000
GAT Asp	TCT Ser	GGT Gly	GTT Val	ATG Met 200	Lys	GTA Val	GGT Gly	TAC	ACT Thr 201	Glu	ATA Ile	GAT Asp	GGC Gly	AAG Lys 201	His	6048
TTC Phe	TAC	TTT Phe	GCT Ala 202	Glu	AAC Asn	GGA Gly	GAA Glu	ATG Met 202	Gln	ATA Ile	GGA Gly	GTA Val	TTT Phe 203	Asn	ACA Thr	6096
GAA Glu	GAT Asp	GGA Gly 203	Phe	AAA Lys	TAT	TTT Phe	GCT Ala 204	His	CAT	AAT Asn	GAA Glu	GAT Asp 204	Leu	GGA Gly	AAT Asn	6144
GAA Glu	GAA Glu 205	Gly	GAA Glu	GAA Glu	ATC Ile	TCA Ser 205	Tyr	TCT Ser	GGT	ATA Ile	TTA Leu 206	AAT Asn 0	TTC Phe	AAT Asn	AAT Asn	6192
AAA Lys 206	Ile	TAC Tyr	TAT Tyr	TTT Phe	GAT Asp 2070	Asp	TCA Ser	TTT Phe	ACA Thr	GCT Ala 2075	Val	GTT Val	GGA Gly	TGG Trp	AAA Lys 2080	6240
GAT Asp	TTA Leu	GAG Glu	GAT Asp	GGT Gly 2089	Ser	AAG Lys	TAT Tyr	TAT Tyr	TTT Phe 2090	Asp	GAA Glu	GAT Asp	ACA Thr	GCA Ala 2095	Glu	6288
GCA Ala	TAT Tyr	ATA Ile	GGT Gly 2100	Leu	TCA Ser	TTA Leu	ATA Ile	AAT Asn 2109	Asp	GGT Gly	CAA Gln	TAT Tyr	TAT Tyr 2110	Phe	AAT Asn	6336
GAT Asp	GAT Asp	GGA Gly 2115	Ile	ATG Met	CAA Gln	GTT Val	GGA Gly 2120	Phe	GTC Val	ACT Thr	ATA Ile	AAT Asn 2125	Asp	AAA Lys	GTC Val	6384
TTC Phe	TAC Tyr 213	Phe	TCT Ser	GAC Asp	TCT Ser	GGA Gly 2135	Ile	ATA. Ile	GAA Glu	TCT Ser	GGA Gly 2140	GTA Val	CAA Gln	AAC Asn	ATA Ile	6432
GAT Asp 2145	Asp	AAT Asn	TAT Tyr	TTC Phe	TAT Tyr 2150	Ile	GAT Asp	GAT Asp	AAT Asn	GGT Gly 2155	Ile	GTT Val	CAA Gln	Ile	GGT Gly 2160	6480
GTA Val	TTT Phe	GAT Asp	Thr	TCA Ser 2165	Asp	GGA Gly	TAT Tyr	AAA Lys	TAT Tyr 2170	Phe	GCA Ala	CCT Pro	Ala	AAT Asn 2175	Thr	6528
GTA Val	AAT Asn	GAT Asp	AAT Asn 2180	Ile	TAC Tyr	GGA Gly	Gln	GCA Ala 2185	Val	GAA Glu	TAT Tyr	AGT Ser	GGT Gly 2190	Leu	GTT Val	6576
AGA Arg	GTT Val	GGG Gly 2195	Glu	GAT Asp	GTA Val	Tyr	TAT Tyr 2200	Phe	GGA Gly	GAA Glu	ACA Thr	TAT Tyr 2205	Thr	ATT (GAG Glu	6624

ACT Thr	GGA Gly 221	TTP	ATA Ile	TAT	GAT Asp	Met 221	Glu	AAT Asn	GAA Glu	AGT Ser	GAT Asp 222	Lys	TAT	TAT	TTC Phe		6672
AAT Asn 222	PLO	GAA Glu	ACT Thr	AAA Lys	AAA Lys 223	Ala	TGC Cys	AAA Lys	GGT Gly	ATT Ile 223	Asn	TTA Leu	ATT	GAT Asp	GAT Asp 2240		6720
ATA Ile	AAA Lys	TAT	TAT Tyr	TTT Phe 224	Asp	GAG Glu	AAG Lys	GGC Gly	ATA Ile 225	Met	AGA Arg	ACG Thr	GGT Gly	CTT Leu 225	Ile		6768
TCA Ser	TTT Phe	GAA Glu	AAT Asn 2260	Asn	AAT Asn	TAT Tyr	TAC Tyr	TTT Phe 226	Asn	GAG Glu	ÀAT Asn	GGT Gly	GAA Glu 227	ATG Met	CAA Gln	1	6816
TTT Phe	GGT Gly	TAT Tyr 227	TTE	AAT Asn	ATA Ile	GAA Glu	GAT Asp 2280	Lys	ATG Met	TTC Phe	TAT Tyr	TTT Phe 228	Gly	GAA Glu	GAT Asp	•	6864
GGT Gly	GTC Val 2290	mec	CAG Gln	ATT Ile	GGA Gly	GTA Val 2299	Phe	AAT Asn	ACA Thr	CCA Pro	GAT Asp 230	Gly	TTT Phe	AAA Lys	TAC Tyr	•	5912
TTT Phe 2305	ALA	CAT His	CAA Gln	AAT Asn	ACT Thr 2310	Leu	GAT Asp	GAG Glu	AAT Asn	TTT Phe 2315	Glu	GGA Gly	GAA Glu	TCA Ser	ATA Ile 2320	6	5960
AAC Asn	TAT Tyr	ACT Thr	GLY	TGG Trp 2325	Leu	GAT Asp	TTA Leu	GAT Asp	GAA Glu 2330	Lys	AGA Arg	TAT Tyr	TAT Tyr	TTT Phe 2335	Thr	7	008
GAT Asp	GAA Glu	TAT Tyr	ATT Ile 2340	Ala	GCA Ala	ACT Thr	GGT Gly	TCA Ser 2345	Val	ATT Ile	ATT Ile	GAT Asp	GGT Gly 2350	GAG Glu	GAG Glu	7	056
TAT Tyr	Tyr	TTT Phe 2355	GAT Asp	CCT Pro	GAT Asp	Thr	GCT Ala 2360	Gln :	TTA Leu	GTG Val	ATT Ile	AGT Ser 2365	Glu			7	098
TAG		 -														7	101

(2) INFORMATION FOR SEQ ID NO:10:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 2366 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:10:

Met Ser Leu Val Asn Arg Lys Gln Leu Glu Lys Met Ala Asn Val Arg 1 5 10 15

Phe Arg Thr Gln Glu Asp Glu Tyr Val Ala Ile Leu Asp Ala Leu Glu 20 25 30

Glu Tyr His Asn Met Ser Glu Asn Thr Val Val Glu Lys Tyr Leu Lys
35 40 45

- Leu Lys Asp Ile Asn Ser Leu Thr Asp Ile Tyr Ile Asp Thr Tyr Lys
 50 60
- Lys Ser Gly Arg Asn Lys Ala Leu Lys Lys Phe Lys Glu Tyr Leu Val 65 70 75 80
- Thr Glu Val Leu Glu Leu Lys Asn Asn Leu Thr Pro Val Glu Lys
 85 90 95
- Asn Leu His Phe Val Trp Ile Gly Gly Gln Ile Asn Asp Thr Ala Ile 100 105 110
- Asn Tyr Ile Asn Gln Trp Lys Asp Val Asn Ser Asp Tyr Asn Val Asn 115 120 125
- Val Phe Tyr Asp Ser Asn Ala Phe Leu Ile Asn Thr Leu Lys Lys Thr 130 135 140
- Val Val Glu Ser Ala Ile Asn Asp Thr Leu Glu Ser Phe Arg Glu Asn 145 150 155 160
- Leu Asn Asp Pro Arg Phe Asp Tyr Asn Lys Phe Phe Arg Lys Arg Met
 165 170 175
- Glu Ile Ile Tyr Asp Lys Gln Lys Asn Phe Ile Asn Tyr Tyr Lys Ala 180 185 190
- Gln Arg Glu Glu Asn Pro Glu Leu Ile Ile Asp Asp Ile Val Lys Thr 195 200 205
- Tyr Leu Ser Asn Glu Tyr Ser Lys Glu Ile Asp Glu Leu Asn Thr Tyr 210 215 220
- Ile Glu Glu Ser Leu Asn Lys Ile Thr Gln Asn Ser Gly Asn Asp Val 225 230 235 240
- Arg Asn Phe Glu Glu Phe Lys Asn Gly Glu Ser Phe Asn Leu Tyr Glu 245 250 255
- Gln Glu Leu Val Glu Arg Trp Asn Leu Ala Ala Ala Ser Asp Ile Leu 260 265 270
- Arg Ile Ser Ala Leu Lys Glu Ile Gly Gly Met Tyr Leu Asp Val Asp 275 280 285
- Met Leu Pro Gly Ile Gln Pro Asp Leu Phe Glu Ser Ile Glu Lys Pro 290 295 300
- Ser Ser Val Thr Val Asp Phe Trp Glu Met Thr Lys Leu Glu Ala Ile 305 310 315 320
- Met Lys Tyr Lys Glu Tyr Ile Pro Glu Tyr Thr Ser Glu His Phe Asp 325 330 335
- Met Leu Asp Glu Glu Val Gln Ser Ser Phe Glu Ser Val Leu Ala Ser 340 345 350
- Lys Ser Asp Lys Ser Glu Ile Phe Ser Ser Leu Gly Asp Met Glu Ala 355 360 365
- Ser Pro Leu Glu Val Lys Ile Ala Phe Asn Ser Lys Gly Ile Ile Asn 370 375 380

- Gln Gly Leu Ile Ser Val Lys Asp Ser Tyr Cys Ser Asn Leu Ile Val 385 390 395 400
- Lys Gln Ile Glu Asn Arg Tyr Lys Ile Leu Asn Asn Ser Leu Asn Pro
 405 410 415
- Ala Ile Ser Glu Asp Asn Asp Phe Asn Thr Thr Thr Asn Thr Phe Ile
 420 425 430
- Asp Ser Ile Met Ala Glu Ala Asn Ala Asp Asn Gly Arg Phe Met Met 435
- Glu Leu Gly Lys Tyr Leu Arg Val Gly Phe Phe Pro Asp Val Lys Thr 450 455 460
- Thr Ile Asn Leu Ser Gly Pro Glu Ala Tyr Ala Ala Ala Tyr Gln Asp 470 475 480
- Leu Leu Met Phe Lys Glu Gly Ser Met Asn Ile His Leu Ile Glu Ala 485 490 495
- Asp Leu Arg Asn Phe Glu Ile Ser Lys Thr Asn Ile Ser Gln Ser Thr 500 505 510
- Glu Gln Glu Met Ala Ser Leu Trp Ser Phe Asp Asp Ala Arg Ala Lys
 515 520 525
- Ala Gln Phe Glu Glu Tyr Lys Arg Asn Tyr Phe Glu Gly Ser Leu Gly 530 535
- Glu Asp Asp Asn Leu Asp Phe Ser Gln Asn Ile Val Val Asp Lys Glu
 545 550 555 560
- Tyr Leu Leu Glu Lys Ile Ser Ser Leu Ala Arg Ser Ser Glu Arg Gly
 565 570 575
- Tyr Ile His Tyr Ile Val Gln Leu Gln Gly Asp Lys Ile Ser Tyr Glu 580 585 590
- Ala Ala Cys Asn Leu Phe Ala Lys Thr Pro Tyr Asp Ser Val Leu Phe 595 600 605
- Gln Lys Asn Ile Glu Asp Ser Glu Ile Ala Tyr Tyr Tyr Asn Pro Gly 610 615 620
- Asp Gly Glu Ile Gln Glu Ile Asp Lys Tyr Lys Ile Pro Ser Ile Ile 625 630 635 640
- Ser Asp Arg Pro Lys Ile Lys Leu Thr Phe Ile Gly His Gly Lys Asp 645 650 655
- Glu Phe Asn Thr Asp Ile Phe Ala Gly Phe Asp Val Asp Ser Leu Ser 660 665 670
- Thr Glu Ile Glu Ala Ala Ile Asp Leu Ala Lys Glu Asp Ile Ser Pro 675 680 685
- Lys Ser Ile Glu Ile Asn Leu Leu Gly Cys Asn Met Phe Ser Tyr Ser 690 695 700
- Ile Asn Val Glu Glu Thr Tyr Pro Gly Lys Leu Leu Leu Lys Val Lys 710 715 720

- Asp Lys Ile Ser Glu Leu Met Pro Ser Ile Ser Gln Asp Ser Ile Ile 725 730 735
- Val Ser Ala Asn Gln Tyr Glu Val Arg Ile Asn Ser Glu Gly Arg Arg 740 745 750
- Glu Leu Leu Asp His Ser Gly Glu Trp Ile Asn Lys Glu Glu Ser Ile 755 760 765
- Ile Lys Asp Ile Ser Ser Lys Glu Tyr Ile Ser Phe Asn Pro Lys Glu
 770 780
- Asn Lys Ile Thr Val Lys Ser Lys Asn Leu Pro Glu Leu Ser Thr Leu 785 790 795 800
- Leu Gln Glu Ile Arg Asn Asn Ser Asn Ser Ser Asp Ile Glu Leu Glu 805 810 815
- Glu Lys Val Met Leu Thr Glu Cys Glu Ile Asn Val Ile Ser Asn Ile 820 825 830
- Asp Thr Gln Ile Val Glu Glu Arg Ile Glu Glu Ala Lys Asn Leu Thr 835 840 845
- Ser Asp Ser Ile Asn Tyr Ile Lys Asp Glu Phe Lys Leu Ile Glu Ser 850 855 860
- Ile Ser Asp Ala Leu Cys Asp Leu Lys Gln Gln Asn Glu Leu Glu Asp 865 870 875 880
- Ser His Phe Ile Ser Phe Glu Asp Ile Ser Glu Thr Asp Glu Gly Phe 885 890 895
- Ser Ile Arg Phe Ile Asn Lys Glu Thr Gly Glu Ser Ile Phe Val Glu 900 905 910
- Thr Glu Lys Thr Ile Phe Ser Glu Tyr Ala Asn His Ile Thr Glu Glu 915 920 925
- Ile Ser Lys Ile Lys Gly Thr Ile Phe Asp Thr Val Asn Gly Lys Leu 930 935 940
- Val Lys Lys Val Asn Leu Asp Thr Thr His Glu Val Asn Thr Leu Asn 945 955 960
- Ala Ala Phe Phe Ile Gln Ser Leu Ile Glu Tyr Asn Ser Ser Lys Glu 965 970 975
- Ser Leu Ser Asn Leu Ser Val Ala Met Lys Val Gln Val Tyr Ala Gln 980 985 990
- Leu Phe Ser Thr Gly Leu Asn Thr Ile Thr Asp Ala Ala Lys Val Val 995 1000 1005
- Glu Leu Val Ser Thr Ala Leu Asp Glu Thr Ile Asp Leu Leu Pro Thr 1010 1015 1020
- Leu Ser Glu Gly Leu Pro Ile Ile Ala Thr Ile Ile Asp Gly Val Ser 1025 1030 1035 1040
- Leu Gly Ala Ala Ile Lys Glu Leu Ser Glu Thr Ser Asp Pro Leu Leu 1045 1050 1055

- Arg Gln Glu Ile Glu Ala Lys Ile Gly Ile Met Ala Val Asn Leu Thr 1060 1065 1070
- Thr Ala Thr Thr Ala Ile Ile Thr Ser Ser Leu Gly Ile Ala Ser Gly 1075 1080 1085
- Phe Ser Ile Leu Leu Val Pro Leu Ala Gly Ile Ser Ala Gly Ile Pro 1090 1095 1100
- Ser Leu Val Asn Asn Glu Leu Val Leu Arg Asp Lys Ala Thr Lys Val
- Val Asp Tyr Phe Lys His Val Ser Leu Val Glu Thr Glu Gly Val Phe 1125 1130 1135
- Thr Leu Leu Asp Asp Lys Ile Met Met Pro Gln Asp Asp Leu Val Ile 1140 1145 1150
- Ser Glu Ile Asp Phe Asn Asn Asn Ser Ile Val Leu Gly Lys Cys Glu 1155 1160 1165
- Ile Trp Arg Met Glu Gly Gly Ser Gly His Thr Val Thr Asp Asp Ile
 1170 1175 1180
- Asp His Phe Phe Ser Ala Pro Ser Ile Thr Tyr Arg Glu Pro His Leu 1185 1190 1195 1200
- Ser Ile Tyr Asp Val Leu Glu Val Gln Lys Glu Glu Leu Asp Leu Ser 1205 1210 1215
- Lys Asp Leu Met Val Leu Pro Asn Ala Pro Asn Arg Val Phe Ala Trp 1220 1225 1230
- Glu Thr Gly Trp Thr Pro Gly Leu Arg Ser Leu Glu Asn Asp Gly Thr
 1235 1240 1245
- Lys Leu Leu Asp Arg Ile Arg Asp Asn Tyr Glu Gly Glu Phe Tyr Trp 1250 1255 1260
- Arg Tyr Phe Ala Phe Ile Ala Asp Ala Leu Ile Thr Thr Leu Lys Pro 1265 1270 1275 1280
- Arg Tyr Glu Asp Thr Asn Ile Arg Ile Asn Leu Asp Ser Asn Thr Arg 1285 1290 1295
- Ser Phe Ile Val Pro Ile Ile Thr Thr Glu Tyr Ile Arg Glu Lys Leu 1300 1305 1310
- Ser Tyr Ser Phe Tyr Gly Ser Gly Gly Thr Tyr Ala Leu Ser Leu Ser 1315 1320 1325
- Gln Tyr Asn Met Gly Ile Asn Ile Glu Leu Ser Glu Ser Asp Val Trp 1330 1335 1340
- Ile Ile Asp Val Asp Asn Val Val Arg Asp Val Thr Ile Glu Ser Asp 1345 1350 1355 1360
- Lys Ile Lys Lys Gly Asp Leu Ile Glu Gly Ile Leu Ser Thr Leu Ser 1365 1370 1375
- Ile Glu Glu Asn Lys Ile Ile Leu Asn Ser His Glu Ile Asn Phe Ser 1380 1385 1390

- Gly Glu Val Asn Gly Ser Asn Gly Phe Val Ser Leu Thr Phe Ser Ile 1395 1400 1405
- Leu Glu Gly Ile Asn Ala Ile Ile Glu Val Asp Leu Leu Ser Lys Ser 1410 1415 1420
- Tyr Lys Leu Leu Ile Ser Gly Glu Leu Lys Ile Leu Met Leu Asn Ser 1425 1430 1435 1440
- Asn His Ile Gln Gln Lys Ile Asp Tyr Ile Gly Phe Asn Ser Glu Leu 1445 1450 1455
- Gln Lys Asn Ile Pro Tyr Ser Phe Val Asp Ser Glu Gly Lys Glu Asn 1460 1465 1470
- Gly Phe Ile Asn Gly Ser Thr Lys Glu Gly Leu Phe Val Ser Glu Leu 1475 1480 1485
- Pro Asp Val Val Leu Ile Ser Lys Val Tyr Met Asp Asp Ser Lys Pro 1490 1495 1500
- Ser Phe Gly Tyr Tyr Ser Asn Asn Leu Lys Asp Val Lys Val Ile Thr 1505 1510 1515 1520
- Lys Asp Asn Val Asn Ile Leu Thr Gly Tyr Tyr Leu Lys Asp Asp Ile 1525 1530 1535
- Lys Ile Ser Leu Ser Leu Thr Leu Gln Asp Glu Lys Thr Ile Lys Leu 1540 1545 1550
- Asn Ser Val His Leu Asp Glu Ser Gly Val Ala Glu Ile Leu Lys Phe 1555 1560 1565
- Met Asn Arg Lys Gly Asn Thr Asn Thr Ser Asp Ser Leu Met Ser Phe 1570 1580
- Leu Glu Ser Met Asn Ile Lys Ser Ile Phe Val Asn Phe Leu Gln Ser 1585 1590 1595 1600
- Asn Ile Lys Phe Ile Leu Asp Ala Asn Phe Ile Ile Ser Gly Thr Thr 1605 1610 1615
- Ser Ile Gly Gln Phe Glu Phe Ile Cys Asp Glu Asn Asp Asn Ile Gln 1620 1630
- Pro Tyr Phe Ile Lys Phe Asn Thr Leu Glu Thr Asn Tyr Thr Leu Tyr 1635 1640 1645
- Val Gly Asn Arg Gln Asn Met Ile Val Glu Pro Asn Tyr Asp Leu Asp 1650 1655 1660
- Asp Ser Gly Asp Ile Ser Ser Thr Val Ile Asn Phe Ser Gln Lys Tyr 1665 1670 1675 1680
- Leu Tyr Gly Ile Asp Ser Cys Val Asn Lys Val Val Ile Ser Pro Asn 1685 1690 1695
- Ile Tyr Thr Asp Glu Ile Asn Ile Thr Pro Val Tyr Glu Thr Asn Asn 1700 1705 1710
- Thr Tyr Pro Glu Val Ile Val Leu Asp Ala Asn Tyr Ile Asn Glu Lys 1715 1720 1725

- Ile Asn Val Asn Ile Asn Asp Leu Ser Ile Arg Tyr Val Trp Ser Asn 1730 1735 1740
- Asp Gly Asn Asp Phe Ile Leu Met Ser Thr Ser Glu Glu Asn Lys Val 1745 1750 1755 1760
- Ser Gln Val Lys Ile Arg Phe Val Asn Val Phe Lys Asp Lys Thr Leu 1765 1770 1775
- Ala Asn Lys Leu Ser Phe Asn Phe Ser Asp Lys Gln Asp Val Pro Val 1780 1785 1790
- Ser Glu Ile Ile Leu Ser Phe Thr Pro Ser Tyr Tyr Glu Asp Gly Leu 1795 1800 1805
- Ile Gly Tyr Asp Leu Gly Leu Val Ser Leu Tyr Asn Glu Lys Phe Tyr 1810 1815 1820
- Ile Asn Asn Phe Gly Met Met Val Ser Gly Leu Ile Tyr Ile Asn Asp 1825 1830 1835 1840
- Ser Leu Tyr Tyr Phe Lys Pro Pro Val Asn Asn Leu Ile Thr Gly Phe 1845 1850 1855
- Val Thr Val Gly Asp Asp Lys Tyr Tyr Phe Asn Pro Ile Asn Gly Gly
 1860 1865 1870
- Ala Ala Ser Ile Gly Glu Thr Ile Ile Asp Asp Lys Asn Tyr Tyr Phe 1875 1880 1885
- Asn Gln Ser Gly Val Leu Gln Thr Gly Val Phe Ser Thr Glu Asp Gly 1890 1895 1900
- Phe Lys Tyr Phe Ala Pro Ala Asn Thr Leu Asp Glu Asn Leu Glu Gly 1905 1910 1915 1920
- Glu Ala Ile Asp Phe Thr Gly Lys Leu Ile Ile Asp Glu Asn Ile Tyr 1925 1930 1935
- Tyr Phe Asp Asp Asn Tyr Arg Gly Ala Val Glu Trp Lys Glu Leu Asp 1940 1950
- Gly Glu Met His Tyr Phe Ser Pro Glu Thr Gly Lys Ala Phe Lys Gly 1955 1960 1965
- Leu Asn Gln Ile Gly Asp Tyr Lys Tyr Tyr Phe Asn Ser Asp Gly Val 1970 1975 1980
- Met Gln Lys Gly Phe Val Ser Ile Asn Asp Asn Lys His Tyr Phe Asp 1985 1990 1995 2000
- Asp Ser Gly Val Met Lys Val Gly Tyr Thr Glu Ile Asp Gly Lys His 2005 2010 2015
- Phe Tyr Phe Ala Glu Asn Gly Glu Met Gln Ile Gly Val Phe Asn Thr 2020 2025 2030
- Glu Asp Gly Phe Lys Tyr Phe Ala His His Asn Glu Asp Leu Gly Asn 2035 2040 2045
- Glu Glu Glu Glu Ile Ser Tyr Ser Gly Ile Leu Asn Phe Asn Asn 2050 2055 2060

- Lys Ile Tyr Tyr Phe Asp Asp Ser Phe Thr Ala Val Val Gly Trp Lys 2065 2070 2075 2080
- Asp Leu Glu Asp Gly Ser Lys Tyr Tyr Phe Asp Glu Asp Thr Ala Glu 2085 2090 2095
- · Ala Tyr Ile Gly Leu Ser Leu Ile Asn Asp Gly Gln Tyr Tyr Phe Asn 2100 2105 2110
 - Asp Asp Gly Ile Met Gln Val Gly Phe Val Thr Ile Asn Asp Lys Val 2115 2120 2125
- Phe Tyr Phe Ser Asp Ser Gly Ile Ile Glu Ser Gly Val Gln Asn Ile 2130 2135 2140
- Asp Asp Asn Tyr Phe Tyr Ile Asp Asp Asn Gly Ile Val Gln Ile Gly 2145 2150 2155 2160
- Val Phe Asp Thr Ser Asp Gly Tyr Lys Tyr Phe Ala Pro Ala Asn Thr 2165 2170 2175
- Val Asn Asp Asn Ile Tyr Gly Gln Ala Val Glu Tyr Ser Gly Leu Val 2180 2185 2190
- Arg Val Gly Glu Asp Val Tyr Tyr Phe Gly Glu Thr Tyr Thr Ile Glu 2195 2200 2205
- Thr Gly Trp Ile Tyr Asp Met Glu Asn Glu Ser Asp Lys Tyr Tyr Phe 2210 2215 2220
- Asn Pro Glu Thr Lys Lys Ala Cys Lys Gly Ile Asn Leu Ile Asp Asp 2225 2230 2235 2240
- Ile Lys Tyr Tyr Phe Asp Glu Lys Gly Ile Met Arg Thr Gly Leu Ile 2245 2250 2255
- Ser Phe Glu Asn Asn Asn Tyr Tyr Phe Asn Glu Asn Gly Glu Met Gln 2260 2270
- Phe Gly Tyr Ile Asn Ile Glu Asp Lys Met Phe Tyr Phe Gly Glu Asp 2275 2280 2285
- Gly Val Met Gln Ile Gly Val Phe Asn Thr Pro Asp Gly Phe Lys Tyr 2290 2295 2300
- Phe Ala His Gln Asn Thr Leu Asp Glu Asn Phe Glu Gly Glu Ser Ile 2305 2310 2315 2320
- Asn Tyr Thr Gly Trp Leu Asp Leu Asp Glu Lys Arg Tyr Tyr Phe Thr 2325 2330 2335
- Asp Glu Tyr Ile Ala Ala Thr Gly Ser Val Ile Ile Asp Gly Glu Glu 2340 2345 2350
- Tyr Tyr Phe Asp Pro Asp Thr Ala Gln Leu Val Ile Ser Glu 2355 2360 2365

(2)	INFO	RMATION FOR SEQ ID NO:11:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 19 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:11:	
TAGA	LAAAA	AT GGCAAATGT	19
(2)	INFO	RMATION FOR SEQ ID NO:12:	
	(i) _.	SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:12:	
TTTC	ATCT	rg tagagtcaaa g	21
(2)	INFO	RMATION FOR SEQ ID NO:13:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:13:	
GATG	CCAC	AA GATGATTTAG TG	22
(2)	INFOR	RMATION FOR SEQ ID NO:14:	
	(i)	SEQUENCE CHARACTERISTICS: (A) LENGTH: 22 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	'n
	(ii)	MOLECULE TYPE: DNA (genomic)	
	(xi)	SEQUENCE DESCRIPTION: SEQ ID NO:14:	
CTAA	TTGAC	GC TGTATCAGGA TC	22

	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:15:	
CGGAA	TTCCT AGAAAAATG GCAAATG	27
(2) I	NFORMATION FOR SEQ ID NO:16:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 26 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	•
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:16:	
GCTCT	AGAAT GACCATAAGC TAGCCA	26
(2) I	NFORMATION FOR SEQ ID NO:17:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(ii) MOLECULE TYPE: DNA (genomic)	
(:	(i) SEQUENCE DESCRIPTION: SEQ ID NO:17:	
CGGAA	TTCGA GTTGGTAGAA AGGTGGA	27
(2) I	FORMATION FOR SEQ ID NO:18:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 27 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
(:	i) MOLECULE TYPE: DNA (genomic)	
(:	i) SEQUENCE DESCRIPTION: SEQ ID NO:18:	
CGGAA	TCGG TTATTATCTT AAGGATG	27

(2) INFORMATION FOR SEQ ID NO:15:

- (2) INFORMATION FOR SEQ ID NO:19:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 28 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:19:

CGGAATTCTT GATAACTGGA TTTGTGAC

(2) INFORMATION FOR SEQ ID NO:20:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 511 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:20:

Leu Ile Thr Gly Phe Val Thr Val Gly Asp Asp Lys Tyr Tyr Phe Asn 1 5 10 15

28

Pro Ile Asn Gly Gly Ala Ala Ser Ile Gly Glu Thr Ile Ile Asp Asp 20 25 30

Lys Asn Tyr Tyr Phe Asn Gln Ser Gly Val Leu Gln Thr Gly Val Phe 35 40 45

Ser Thr Glu Asp Gly Phe Lys Tyr Phe Ala Pro Ala Asn Thr Leu Asp 50 55

Glu Asn Leu Glu Gly Glu Ala Ile Asp Phe Thr Gly Lys Leu Ile Ile 65 70 75 80

Asp Glu Asn Ile Tyr Tyr Phe Asp Asp Asn Tyr Arg Gly Ala Val Glu 85 90 95

Trp Lys Glu Leu Asp Gly Glu Met His Tyr Phe Ser Pro Glu Thr Gly
100 105 110

Lys Ala Phe Lys Gly Leu Asn Gln Ile Gly Asp Tyr Lys Tyr Tyr Phe
115 120 125

Asn Ser Asp Gly Val Met Gln Lys Gly Phe Val Ser Ile Asn Asp Asn 130 135 140

Lys His Tyr Phe Asp Asp Ser Gly Val Met Lys Val Gly Tyr Thr Glu 145 150 155 160

Ile Asp Gly Lys His Phe Tyr Phe Ala Glu Asn Gly Glu Met Gln Ile 165 170 175

Gly Val Phe Asn Thr Glu Asp Gly Phe Lys Tyr Phe Ala His His Asn 180 185 190 Glu Asp Leu Gly Asn Glu Glu Gly Glu Glu Ile Ser Tyr Ser Gly Ile
195 200 205

Leu Asn Phe Asn Asn Lys Ile Tyr Tyr Phe Asp Asp Ser Phe Thr Ala 210 215 220

Val Val Gly Trp Lys Asp Leu Glu Asp Gly Ser Lys Tyr Tyr Phe Asp 225 230 235 240

Glu Asp Thr Ala Glu Ala Tyr Ile Gly Leu Ser Leu Ile Asn Asp Gly 245 250 255

Gln Tyr Tyr Phe Asn Asp Asp Gly Ile Met Gln Val Gly Phe Val Thr 260 265 270

Ile Asn Asp Lys Val Phe Tyr Phe Ser Asp Ser Gly Ile Ile Glu Ser 275 280 285

Gly Val Gln Asn Ile Asp Asp Asn Tyr Phe Tyr Ile Asp Asp Asn Gly
290 295 300

Ile Val Gln Ile Gly Val Phe Asp Thr Ser Asp Gly Tyr Lys Tyr Phe 305 310 315 320

Ala Pro Ala Asn Thr Val Asn Asp Asn Ile Tyr Gly Gln Ala Val Glu 325 330 335

Tyr Ser Gly Leu Val Arg Val Gly Glu Asp Val Tyr Tyr Phe Gly Glu 340 345 350

Thr Tyr Thr Ile Glu Thr Gly Trp Ile Tyr Asp Met Glu Asn Glu Ser 355 360 365

Asp Lys Tyr Tyr Phe Asn Pro Glu Thr Lys Lys Ala Cys Lys Gly Ile 370 375 380

Asn Leu Ile Asp Asp Ile Lys Tyr Tyr Phe Asp Glu Lys Gly Ile Met 385 390 395 400

Arg Thr Gly Leu Ile Ser Phe Glu Asn Asn Asn Tyr Tyr Phe Asn Glu
405 410 415

Asn Gly Glu Met Gln Phe Gly Tyr Ile Asn Ile Glu Asp Lys Met Phe 420 425 430

Tyr Phe Gly Glu Asp Gly Val Met Gln Ile Gly Val Phe Asn Thr Pro 435 440 445

Asp Gly Phe Lys Tyr Phe Ala His Gln Asn Thr Leu Asp Glu Asn Phe 450 450 460

Glu Gly Glu Ser Ile Asn Tyr Thr Gly Trp Leu Asp Leu Asp Glu Lys 465 470 475 480

Arg Tyr Tyr Phe Thr Asp Glu Tyr Ile Ala Ala Thr Gly Ser Val Ile 485 490 495

Ile Asp Gly Glu Glu Tyr Tyr Phe Asp Pro Asp Thr Ala Gln Leu 500 505 510

(2) INFORMATION FOR SEQ ID NO:21:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 608 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: unknown
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:21:

Ser Glu Glu Asn Lys Val Ser Gln Val Lys Ile Arg Phe Val Asn Val

Phe Lys Asp Lys Thr Leu Ala Asn Lys Leu Ser Phe Asn Phe Ser Asp 20 25 30

Lys Gln Asp Val Pro Val Ser Glu Ile Ile Leu Ser Phe Thr Pro Ser
35 40 45

Tyr Tyr Glu Asp Gly Leu Ile Gly Tyr Asp Leu Gly Leu Val Ser Leu 50 55 60

Tyr Asn Glu Lys Phe Tyr Ile Asn Asn Phe Gly Met Met Val Ser Gly 65 70 75 80

Leu Ile Tyr Ile Asn Asp Ser Leu Tyr Tyr Phe Lys Pro Pro Val Asn 85 90 95

Asn Leu Ile Thr Gly Phe Val Thr Val Gly Asp Asp Lys Tyr Tyr Phe
100 105 110

Asn Pro Ile Asn Gly Gly Ala Ala Ser Ile Gly Glu Thr Ile Ile Asp 115 120 125

Asp Lys Asn Tyr Tyr Phe Asn Gln Ser Gly Val Leu Gln Thr Gly Val 130 135 140

Phe Ser Thr Glu Asp Gly Phe Lys Tyr Phe Ala Pro Ala Asn Thr Leu 145 150 155 160

Asp Glu Asn Leu Glu Gly Glu Ala Ile Asp Phe Thr Gly Lys Leu Ile 165 170 175

Ile Asp Glu Asn Ile Tyr Tyr Phe Asp Asp Asn Tyr Arg Gly Ala Val 180 185 190

Glu Trp Lys Glu Leu Asp Gly Glu Met His Tyr Phe Ser Pro Glu Thr 195 200 205

Gly Lys Ala Phe Lys Gly Leu Asn Gln Ile Gly Asp Tyr Lys Tyr Tyr 210 215 220

Phe Asn Ser Asp Gly Val Met Gln Lys Gly Phe Val Ser Ile Asn Asp 225 230 235 240

Asn Lys His Tyr Phe Asp Asp Ser Gly Val Met Lys Val Gly Tyr Thr 245 250 255

Glu Ile Asp Gly Lys His Phe Tyr Phe Ala Glu Asn Gly Glu Met Gln
260 265 270

Ile Gly Val Phe Asn Thr Glu Asp Gly Phe Lys Tyr Phe Ala His His 280 Asn Glu Asp Leu Gly Asn Glu Glu Glu Glu Glu Ile Ser Tyr Ser Gly Ile Leu Asn Phe Asn Asn Lys Ile Tyr Tyr Phe Asp Asp Ser Phe Thr 310 Ala Val Val Gly Trp Lys Asp Leu Glu Asp Gly Ser Lys Tyr Tyr Phe Asp Glu Asp Thr Ala Glu Ala Tyr Ile Gly Leu Ser Leu Ile Asn Asp Gly Gln Tyr Tyr Phe Asn Asp Asp Gly Ile Met Gln Val Gly Phe Val 360 Thr Ile Asn Asp Lys Val Phe Tyr Phe Ser Asp Ser Gly Ile Ile Glu Ser Gly Val Gln Asn Ile Asp Asp Asn Tyr Phe Tyr Ile Asp Asp Asn 390 Gly Ile Val Gln Ile Gly Val Phe Asp Thr Ser Asp Gly Tyr Lys Tyr 405 410 Phe Ala Pro Ala Asn Thr Val Asn Asp Asn Ile Tyr Gly Gln Ala Val 425 Glu Tyr Ser Gly Leu Val Arg Val Gly Glu Asp Val Tyr Tyr Phe Gly Glu Thr Tyr Thr Ile Glu Thr Gly Trp Ile Tyr Asp Met Glu Asn Glu Ser Asp Lys Tyr Tyr Phe Asn Pro Glu Thr Lys Lys Ala Cys Lys Gly 475 Ile Asn Leu Ile Asp Asp Ile Lys Tyr Tyr Phe Asp Glu Lys Gly Ile 490 Met Arg Thr Gly Leu Ile Ser Phe Glu Asn Asn Asn Tyr Tyr Phe Asn Glu Asn Gly Glu Met Gln Phe Gly Tyr Ile Asn Ile Glu Asp Lys Met 520

Phe Tyr Phe Gly Glu Asp Gly Val Met Gln Ile Gly Val Phe Asn Thr 530 535 540

Pro Asp Gly Phe Lys Tyr Phe Ala His Gln Asn Thr Leu Asp Glu Asn 545 550 555 560

Phe Glu Gly Glu Ser Ile Asn Tyr Thr Gly Trp Leu Asp Leu Asp Glu 565 570 575

Lys Arg Tyr Tyr Phe Thr Asp Glu Tyr Ile Ala Ala Thr Gly Ser Val

Ile Ile Asp Gly Glu Glu Tyr Tyr Phe Asp Pro Asp Thr Ala Gln Leu 595 600 605

	(:		(A) : (B) : (C) :	LENG' Type Strai	TH: : nu NDEDI	ACTE 1330 cleic NESS : lir	base c ac: : do:	e pa id	irs							
	(ii	L) MO	OLECT	OTE 1	YPE	: DNA	4 (ge	enom:	ic)							
	(ix	1		NAME/		CDS		L								
	(xi	.) SE	EQUEN	VCE I	ESCF	RIPTI	ON:	SEQ	ID N	10:22	:					
ATG Met	. ALA	CGT Arg	CTC Lev	CTG Leu 5	Ser	ACC Thr	TTC Phe	ACT Thr	GAA Glu	Tyr	ATC Ile	AAG Lys	AA(Asr	ATC Ile	ATC Ile	4
AAT Asn	ACC Thr	TCC	ATC Ile 20	: Leu	AAC Asn	CTG Leu	CGC	TAC Tyr 25	Glu	TCC Ser	AAT Asn	CAC His	CTC Leu	ı Ile	GAC Asp	9
CŤG Leu	TCT Ser	CGC Arg 35	Tyr	GCT Ala	TCC Ser	AAA Lys	ATC Ile 40	Asn	ATC Ile	GGT Gly	TCT Ser	AAA Lys 45	Val	' AAC Asn	TTC	14
GAT Asp	CCG Pro 50	ATC Ile	GAC Asp	AAG Lys	AAT Asn	CAG Gln 55	ATC Ile	CAG Gln	CTG Leu	TTC Phe	AAT Asn 60	Leu	GAA Glu	TCT Ser	TCC Ser	192
AAA Lys 65	ATC Ile	GAA Glu	GTT Val	ATC Ile	CTG Leu 70	AAG Lys	AAT Asn	GCT Ala	ATC Ile	GTA Val 75	TAC Tyr	AAC Asn	TCT Ser	ATG Met	TAC Tyr 80	240
GAA Glu	AAC Asn	TTC Phe	TCC	ACC Thr 85	TCC Ser	TTC Phe	TGG Trp	ATC Ile	CGT Arg 90	ATC Ile	CCG Pro	AAA Lys	TAC Tyr	TTC Phe 95	AAC Asn	288
TCC Ser	ATC	TCT Ser	CTG Leu 100	AAC Asn	AAT Asn	GAA Glu	TAC Tyr	ACC Thr 105	ATC Ile	ATC Ile	AAC Asn	TGC Cys	ATG Met 110	GAA Glu	AAC Asn	336
AAT Asn	TCT Ser	GGT Gly 115	TGG Trp	AAA Lys	GTA Val	TCT Ser	CTG Leu 120	AAC Asn	TAC Tyr	.GGT Gly	GAA Glu	ATC Ile 125	ATC Ile	TGG Trp	ACT Thr	384
CTG Leu	CAG Gln 130	GAC Asp	ACT Thr	CAG Gln	GAA Glu	ATC Ile 135	AAA Lys	CAG Gln	CGT Arg	GTT Val	GTA Val 140	TTC Phe	AAA Lys	TAC Tyr	TCT Ser	432
CAG Gln L45	ATG Met	ATC Ile	AAC Asn	ATC Ile	TCT Ser 150	GAC Asp	TAC Tyr	ATC Ile	AAT Asn	CGC Arg 155	TGG Trp	ATC Ile	TTC Phe	GTT Val	ACC Thr 160	480
ATC le	ACC Thr	AAC Asn	AAT Asn	CGT Arg 165	CTG Leu	AAT Asn	AAC Asn	TCC Ser	AAA Lys 170	ATC Ile	TAC Tyr	ATC Ile	AAC Asn	GGC Gly 175	CGT Arg	528

(2) INFORMATION FOR SEQ ID NO:22:

CT(Le	3 ATO	C GAG	CAG Gln 180	Lys	A CCC	ATC Ile	TCC Ser	AA: Asi 185	ı Lev	GGT Gly	AA(Asr	ATO	CA(His	Ala	TCT Ser	576
AA1 Asr	AAC Asr	TATO 11e	e met	Phe	AAA Lys	CTG Leu	GAC Asp 200	Gl	TGI Cys	CGT Arg	GAC Asp	Thr 205	His	C CGC	TAC Tyr	624
ATC Ile	TGG Trp 210	TTE	AAA Lys	TAC	TTC Phe	AAT Asn 215	CTG Leu	TTC	GAC Asp	Lys	GAA Glu 220	Leu	AAC Asn	GAA Glu	AAA Lys	672
GAA Glu 225	тте	AAA Lys	GAC Asp	CTG Leu	TAC Tyr 230	Asp	AAC Asn	CAG Gln	TCC Ser	AAT Asn 235	Ser	GGT	ATC Ile	CTG Leu	AAA Lys 240	720
GAC Asp	TTC Phe	TGG	GGT Gly	GAC Asp 245	TAC	CTG Leu	CAG Gln	TAC	GAC Asp 250	AAA Lys	CCG Pro	TAC	TAC	ATG Met 255	CTG Leu	768
AAT Asn	CTG Leu	TAC Tyr	GAT Asp 260	CCG Pro	AAC Asn	AAA Lys	TAC	GTT Val 265	GAC Asp	GTC Val	AAC Asn	AAT Asn	GTA Val 270	GGT Gly	ATC Ile	816
CGC Arg	GGT Gly	TAC Tyr 275	ATG Met	TAC Tyr	CTG Leu	AAA Lys	GGT Gly 280	CCG Pro	CGT Arg	GGT Gly	TCT Ser	GTT Val 285	ATG Met	ACT Thr	ACC Thr	864
AAC Asn	ATC Ile 290	TAC Tyr	CTG Leu	AAC Asn	TCT Ser	TCC Ser 295	CTG Leu	TAC Tyr	CGT Arg	GGT Gly	ACC Thr 300	AAA Lys	TTC Phe	ATC Ile	ATC Ile	912
AAG Lys 305	AAA Lys	TAC Tyr	GCG Ala	TCT Ser	GGT Gly 310	AAC Asn	AAG Lys	GAC Asp	AAT Asn	ATC Ile 315	GTT Val	CGC Ar ģ	AAC Asn	AAT Asn	GAT Asp 320	960
CGT	GTA Val	TAC Tyr	ATC Ile	AAT Asn 325	GTT Val	GTA Val	GTT Val	AAG Lys	AAC Asn 330	AAA Lys	GAA Glu	TAC Tyr	CGT Arg	CTG Leu 335	GCT Ala	1008
ACC Thr	AAT Asn	GCT Ala	TCT Ser 340	CAG Gln	GCT Ala	GGT Gly	Val	GAA Glu 345	AAG Lys	ATC Ile	TTG Leu	TCT Ser	GCT Ala 350	CTG Leu	GAA Glu	1056
ATC Ile	CCG Pro	GAC Asp 355	GTT Val	GGT Gly	AAT Asn	Leu	TCT Ser 360	CAG Gln	GTA Val	GTT Val	GTA Val	ATG Met 365	AAA Lys	TCC Ser	AAG Lys	1104
AAC Asn	GAC Asp 370	CAG Gln	GGT Gly	ATC Ile	Thr	AAC Asn 375	AAA ' Lys '	TGC Cys	AAA Lys	Met	AAT Asn 380	CTG Leu	CAG Gln	GAC Asp	AAC Asn	1152
AAT Asn 385	GGT Gly	AAC Asn	GAT . Asp	Ile	GGT Gly 390	TTC . Phe	ATC (GGT Gly	Phe :	CAC His 395	CAG Gln	TTC Phe	AAC Asn	AAT Asn	ATC Ile 400	1200
GCT Ala	AAA Lys	CTG Leu	Val .	GCT Ala 405	TCC . Ser .	AAC ' Asn '	IGG (Гуr	AAT Asn 2 410	CGT (Arg (CAG . Gln	ATC Ile	Glu	CGT Arg 415	TCC Ser	1248

TCT CGC ACT CTG GGT TGC TCT TGG GAG TTC ATC CCG GTT GAT GAC GGT
Ser Arg Thr Leu Gly Cys Ser Trp Glu Phe Ile Pro Val Asp Asp Gly
420
425
430

TGG GGT GAA CGT CCG CTG TAACCCGGGA AAGCTT
Trp Gly Glu Arg Pro Leu
435

1330

(2) INFORMATION FOR SEQ ID NO:23:

(i) SEQUENCE CHARACTERISTICS:

- (A) LENGTH: 438 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:23:

Met Ala Arg Leu Leu Ser Thr Phe Thr Glu Tyr Ile Lys Asn Ile Ile
1 5 10 15

Asn Thr Ser Ile Leu Asn Leu Arg Tyr Glu Ser Asn His Leu Ile Asp
20 25 30

Leu Ser Arg Tyr Ala Ser Lys Ile Asn Ile Gly Ser Lys Val Asn Phe
35 40 45

Asp Pro Ile Asp Lys Asn Gln Ile Gln Leu Phe Asn Leu Glu Ser Ser 50 55 60

Lys Ile Glu Val Ile Leu Lys Asn Ala Ile Val Tyr Asn Ser Met Tyr 65 70 75 80

Glu Asn Phe Ser Thr Ser Phe Trp Ile Arg Ile Pro Lys Tyr Phe Asn 85 90 95

Ser Ile Ser Leu Asn Asn Glu Tyr Thr Ile Ile Asn Cys Met Glu Asn 100 105 110

Asn Ser Gly Trp Lys Val Ser Leu Asn Tyr Gly Glu Ile Ile Trp Thr 115 120 125

Leu Gln Asp Thr Gln Glu Ile Lys Gln Arg Val Val Phe Lys Tyr Ser 130 135 140

Gln Met Ile Asn Ile Ser Asp Tyr Ile Asn Arg Trp Ile Phe Val Thr 145 150 155 160

Ile Thr Asn Asn Arg Leu Asn Asn Ser Lys Ile Tyr Ile Asn Gly Arg 165 170 175

Leu Ile Asp Gln Lys Pro Ile Ser Asn Leu Gly Asn Ile His Ala Ser 180 185 190

Asn Asn Ile Met Phe Lys Leu Asp Gly Cys Arg Asp Thr His Arg Tyr 195 200 205

Ile Trp Ile Lys Tyr Phe Asn Leu Phe Asp Lys Glu Leu Asn Glu Lys 210 215 220 Glu Ile Lys Asp Leu Tyr Asp Asn Gln Ser Asn Ser Gly Ile Leu Lys 225 230 235 240

Asp Phe Trp Gly Asp Tyr Leu Gln Tyr Asp Lys Pro Tyr Tyr Met Leu 245 250 255

Asn Leu Tyr Asp Pro Asn Lys Tyr Val Asp Val Asn Asn Val Gly Ile
260 265 270

Arg Gly Tyr Met Tyr Leu Lys Gly Pro Arg Gly Ser Val Met Thr Thr 275 280 285

Asn Ile Tyr Leu Asn Ser Ser Leu Tyr Arg Gly Thr Lys Phe Ile Ile 290 295 300

Lys Lys Tyr Ala Ser Gly Asn Lys Asp Asn Ile Val Arg Asn Asn Asp 305 310 315 320

Arg Val Tyr Ile Asn Val Val Val Lys Asn Lys Glu Tyr Arg Leu Ala 325 330 335

Thr Asn Ala Ser Gln Ala Gly Val Glu Lys Ile Leu Ser Ala Leu Glu
340 345 350

Ile Pro Asp Val Gly Asn Leu Ser Gln Val Val Wet Lys Ser Lys 355 360 365

Asn Asp Gln Gly Ile Thr Asn Lys Cys Lys Met Asn Leu Gln Asp Asn 370 375 380

Asn Gly Asn Asp Ile Gly Phe Ile Gly Phe His Gln Phe Asn Asn Ile 385 390 395 400

Ala Lys Leu Val Ala Ser Asn Trp Tyr Asn Arg Gln Ile Glu Arg Ser 405 410 415

Ser Arg Thr Leu Gly Cys Ser Trp Glu Phe Ile Pro Val Asp Asp Gly
420 425 430

Trp Gly Glu Arg Pro Leu 435

(2) INFORMATION FOR SEQ ID NO:24:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 23 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: unknown
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:24:

Met Gly His His His His His His His His His Ser Ser Gly His

10
15

Ile Glu Gly Arg His Met Ala
20

	(i	((A) I (B) 1 (C) S	ENGT TYPE : TRAN	TH: 1 nuc TDEDN	ACTER 1402 :leic TESS:	base aci dou	pai .d	irs							
	(ii	.) MC	LECU	TE I	YPE:	DNA	(ge	nomi	.c)·							
	(ix	(AME/		CDS										
	(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:25	:					
ATG Met 1	Gly	CAT	CAT His	CAT His 5	His	CAT His	CAT	CAT His	CAT His 10	CAT His	CAC His	AGC Ser	AGC Ser	GGC Gly 15		48
ATC Ile	GAA Glu	GGT Gly	CGT Arg 20	CAT	ATG Met	GCT Ala	AGC Ser	ATG Met 25	GCT Ala	CGT	CTG Leu	CTG Leu	TCT Ser 30	Thr	TTC Phe	96
ACT Thr	GAA Glu	TAC Tyr 35	ATC Ile	AAG Lys	AAC Asn	ATC Ile	ATC Ile 40	AAT Asn	ACC Thr	TCC Ser	ATC Ile	CTG Leu 45	AAC Asn	CTG Leu	CGC Arg	144
TAC Tyr	GAA Glu 50	Ser	AAT Asn	CAC His	CTG Leu	ATC Ile 55	GAC Asp	CTG Leu	TCT Ser	CGC Arg	TAC Tyr 60	GCT Ala	TCC Ser	AAA Lys	ATC Ile	192
AAC Asn 65	ATC Ile	GGT Gly	TCT Ser	AAA Lys	GTT Val 70	AAC Asn	TTC Phe	GAT Asp	CCG Pro	ATC Ile 75	GAC Asp	AAG Lys	AAT Asn	CAG Gln	ATC Ile 80	240
CAG Gln	CTG Leu	TTC Phe	AAT Asn	CTG Leu 85	GAA Glu	TCT Ser	TCC Ser	AAA Lys	ATC Ile 90	GAA Glu	GTT Val	ATC Ile	CTG Leu	AAG Lys 95	AAT Asn	288
GCT Ala	ATC Ile	GTA Val	TAC Tyr 100	AAC Asn	TCT Ser	ATG Met	TAC Tyr	GAA Glu 105	AAC Asn	TTC Phe	TCC Ser	ACC Thr	TCC Ser 110	TTC Phe	TGG Trp	336
[le	Arg	Ile	Pro	Lys	Tyr	Phe	Asn	Ser	ATC Ile	Ser	Leu	Asn	Asn	GAA Glu	TAC	384
ACC Thr	ATC Ile 130	ATC Ile	AAC Asn	TGC Cys	ATG Met	GAA Glu 135	AAC Asn	AAT Asn	TCT Ser	GGT Gly	TGG Trp 140	AAA Lys	GTA Val	TCT Ser	CTG Leu	432
AC Asn .45	TAC Tyr	GGT Gly	GAA Glu	ATC Ile	ATC Ile 150	TGG Trp	ACT Thr	CTG Leu	CAG Gln	GAC Asp 155	ACT Thr	CAG Gln	GAA Glu	ATC Ile	AAA Lys 160	480
AG In	CGT Arg	GTT Val	GTA Val	TTC Phe 165	AAA Lys	TAC Tyr	TCT Ser	Gln	ATG Met 170	ATC Ile	AAC Asn	ATC Ile	TCT Ser	GAC Asp	TAC Tyr	528

(2) INFORMATION FOR SEQ ID NO:25:

AT Il	C AA e As	T CG n Ar	C TG G Tr 18	בי ע	C TT e Ph	C GT e Va	T AC 1 Th	C AT r Il 18	e Tn	C AA r As	C AA n As	T CG n Ar	T CT g Le 19	u As	T AAC n Asn	576
TC Se	C AA r Ly	A AT s Il 19	e ry	C AT	C AA e As	C GGe n Gl	C CG' y Arg 200	a re	G AT	C GA e As _l	C CA	G AA n Ly: 20!	s Pr	G AT	C TCC e Ser	624
AA' Ası	r CTC n Let 210	a GI	T AAG Y Ası	C AT	C CAG	C GC: s Ala 219	a Sei	r AA'	T AA(C ATO	C ATO	t Phe	Lys	A CTO	G GAC	672
GG1 G1 225	Cy:	r cg	r Gad	C AC	CAC His 230	Arg	TAC Tyr	TATO	C TGC e Tr	3 ATC 5 Ile 235	Lys	A TAC	TTC Phe	AA?	CTG Leu 240	720
TTO	GAC Asp	AA/	A GAA s Glu	Leu 249	l AST	GAA Glu	AAA Lys	GAZ Glu	A ATO 1 Ile 250	: Lys	GAC Asp	C CTG	TAC	GAC Asp 255	AAC Asn	768
CAG Gln	TCC Ser	AA1	TCT Ser 260	GTA	TATO	CTG Leu	AAA Lys	GAC Asp 265	Phe	TGG Trp	GGT Gly	GAC Asp	TAC Tyr 270	Leu	CAG Gln	816
TAC	GAC Asp	Lys 275	PIO	TAC	TAC	ATG Met	CTG Leu 280	AAT Asn	CTG Leu	TAC Tyr	GAT Asp	CCG Pro 285	AAC Asn	AAA Lys	TAC	864
GTT Val	GAC Asp 290	val	AAC Asn	AAT Asn	GTA Val	GGT Gly 295	ATC Ile	CGC Arg	GGT Gly	TAC Tyr	ATG Met 300	TAC Tyr	CTG Leu	AAA Lys	GGT Gly	912
CCG Pro 305	CGT Arg	GGT	TCT	GTT Val	ATG Met 310	ACT Thr	ACC Thr	AAC Asn	ATC Ile	TAC Tyr 315	CTG Leu	AAC Asn	TCT Ser	TCC Ser	CTG Leu 320	960
TAC Tyr	CGT Arg	GGT Gly	ACC Thr	AAA Lys 325	TTC Phe	ATC Ile	ATC Ile	AAG Lys	AAA Lys 330	TAC Tyr	GCG Ala	TCT Ser	GGT Gly	AAC Asn 335	AAG Lys	1008
GAC Asp	AAT Asn	ATC Ile	GTT Val 340	CGC Arg	AAC Asn	AAT Asn	GAT Asp	CGT Arg 345	GTA Val	TAC Tyr	ATC Ile	AAT Asn	GTT Val 350	GTA Val	GTT Val	1056
AAG Lys	AAC Asn	AAA Lys 355	GAA Glu	TAC Tyr	CGT Arg	CTG Leu	GCT Ala 360	ACC Thr	AAT Asn	GCT Ala	TCT Ser	CAG Gln 365	GCT Ala	GGT Gly	GTA Val	1104
GAA Glu	AAG Lys 370	ATC Ile	TTG Leu	TCT Ser	GCT Ala	CTG Leu 375	GAA Glu	ATC Ile	CCG Pro	GAC Asp	GTT Val 380	GGT Gly	AAT Asn	CTG Leu	TCT Ser	1152
CAG Gln 385	GTA Val	GTT Val	GTA Val	ATG Met	AAA Lys 390	TCC Ser	AAG Lys	AAC Asn	GAC Asp	CAG Gln 395	GGT Gly	ATC Ile	ACT Thr	AAC Asn	AAA Lys 400	1200
TGC Cys	AAA Lys	ATG Met	ASN	CTG Leu 405	CAG Gln	GAC Asp	AAC Asn	Asn	GGT Gly 410	AAC Asn	GAT Asp	ATC	Gly	TTC Phe 415	ATC Ile	1248

GG: Gl;	r TTO	C CAG His	C CAC 5 Gl: 420	1 Pue	C AA(C Asi	AA? Asi	T ATO	GC' Ala 42	a Lys	CTC	G GTT 1 Val	r gci L Ala	TC(3 Ser 43(: Asr	TGG Trp	1296
TAC Ty:	AA7 Asr	CGT Arg 435	GTI	ATO	GAZ Glu	A CGT	TCC Ser 440	Se	r CGC	ACT Thr	CTC Lev	GG1 Gly 445	Cys	C TCT	TGG Trp	1344
GAC Glu	TTC Phe 450	: Ile	CCC Pro	GTT Val	GAT Asp	GAC Asp 455	Gly	Tr	GGT Gly	GAA Glu	CGT Arg 460	Pro	CTC Leu	; .		1386
TAA	CCCC	GGA	AAGC	TT												1402
(2)	INF	'ORMA	TION	FOR	SEQ	ID	NO : 2	6:								
	((A (B (D) LE) TY) TO	NGTH PE : POLO	RACT: 46 amin GY: E: p	2 am o ac line	ino id ar	: acid	s						
									0 TD	170	2.0					
								•	Q ID							
Met 1	GIÀ	His	His	His 5	His	His	His	His	His 10	His	His	Ser	Ser	Gly 15	His	
Ile	Glu	Gly	Arg 20	His	Met	Ala	Ser	Met 25	Ala	Arg	Leu	Leu	Ser 30	Thr	Phe	
Thr	Glu	Tyr 35	Ile	Lys	Asn	Ile	Ile 40	Asn	Thr	Ser	Ile	Leu 45	Asn	Leu	Arg	
Tyr	Glu 50	Ser	Asn	His	Leu	Ile 55	Asp	Leu	Ser	Arg	Tyr 60	Ala	Ser	Ĺys	Ile	
Asn 65	Ile	Gly	Ser	Lys	Val 70	Asn	Phe	Asp	Pro	Ile 75	Asp	Lys	Asn	Gln	Ile 80	
Gln	Leu	Phe	Asn	Leu 85	Glu	Ser	Ser	Lys	Ile 90	Glu	Val	Ile	Leu	Lys 95	Asn	
Ala	Ile	Val	Tyr 100	Asn	Ser	Met	Tyr	Glu 105	Asn	Phe	Ser	Thr	Ser 110	Phe	Trp	
Ile	Arg	Ile 115	Pro	Lys	Tyr	Phe	Asn 120	Ser	Ile	Ser	Leu	Asn 125	Àsn	Glu	Tyr	
Thr	Ile 130	Ile	Asn	Cys	Met	Glu 135	Asn	Asn	Ser	Gly	Trp 140	Lys	Val	Ser	Leu	
Asn 145	Tyr	Gly	Glu	Ile	Ile 150	Trp	Thr	Leu	Gln	Asp 155	Thr	Gln	Glu	Ile	Lys 160	
Gln	Arg	Val	Val	Phe 165	Lys	Tyr	Ser	Gln	Met 170	Ile	Asn	Ile	Ser	Asp 175	Tyr	
Ile	Asn	Arg	Trp 180	Ile	Phe	Val	Thr	Ile 185	Thr	Asn	Asn	Arg	Leu 190	Asn	Asn	

Ser Lys Ile Tyr Ile Asn Gly Arg Leu Ile Asp Gln Lys Pro Ile Ser Asn Leu Gly Asn Ile His Ala Ser Asn Asn Ile Met Phe Lys Leu Asp Gly Cys Arg Asp Thr His Arg Tyr Ile Trp Ile Lys Tyr Phe Asn Leu 230 235 Phe Asp Lys Glu Leu Asn Glu Lys Glu Ile Lys Asp Leu Tyr Asp Asn Gln Ser Asn Ser Gly Ile Leu Lys Asp Phe Trp Gly Asp Tyr Leu Gln Tyr Asp Lys Pro Tyr Tyr Met Leu Asn Leu Tyr Asp Pro Asn Lys Tyr Val Asp Val Asn Asn Val Gly Ile Arg Gly Tyr Met Tyr Leu Lys Gly Pro Arg Gly Ser Val Met Thr Thr Asn Ile Tyr Leu Asn Ser Ser Leu Tyr Arg Gly Thr Lys Phe Ile Ile Lys Lys Tyr Ala Ser Gly Asn Lys 325 330 Asp Asn Ile Val Arg Asn Asn Asp Arg Val Tyr Ile Asn Val Val Val Lys Asn Lys Glu Tyr Arg Leu Ala Thr Asn Ala Ser Gln Ala Gly Val Glu Lys Ile Leu Ser Ala Leu Glu Ile Pro Asp Val Gly Asn Leu Ser Gln Val Val Met Lys Ser Lys Asn Asp Gln Gly Ile Thr Asn Lys Cys Lys Met Asn Leu Gln Asp Asn Asn Gly Asn Asp Ile Gly Phe Ile Gly Phe His Gln Phe Asn Asn Ile Ala Lys Leu Val Ala Ser Asn Trp 425 Tyr Asn Arg Gln Ile Glu Arg Ser Ser Arg Thr Leu Gly Cys Ser Trp Glu Phe Ile Pro Val Asp Asp Gly Trp Gly Glu Arg Pro Leu

- (2) INFORMATION FOR SEQ ID NO:27:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 3891 base pairs

455

- (B) TYPE: nucleic acid
- (C) STRANDEDNESS: double
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)

(ix) FEATURE:
(A) NAME/KEY: CDS
(B) LOCATION: 1..3888

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:27:

	_			_					
			CAA Gln					GGT Gly	48
			AAA Lys					CCA Pro	96
			CAT His						144
			GAA Glu 55						192
			TCA Ser						240
			TAT Tyr				-	 	288
			CTT Leu						336
			GGT Gly						384
			ATT Ile 135						432
			CTA Leu						480
			AGC Ser						528
			ACT Thr						576
			TCA Ser						624
			ACA Thr 215						672

CTT Leu 225	Ile	CAT His	GCT Ala	GGA Gly	CAT His 230	AGA Arg	TTA Leu	TAT	GGA Gly	ATA Ile 235	Ala	ATT	AAT Asn	CCA Pro	AAT Asn 240	720
AGG Arg	GTT Val	TTT Phe	AAA Lys	GTA Val 245	AAT Asn	ACT Thr	AAT Asn	GCC Ala	TAT Tyr 250	TAT	GAA Glu	ATG Met	AGT Ser	GGG Gly 255	TTA Leu	768
GAA Glu	GTA Val	AGC Ser	TTT Phe 260	GAG Glu	GAA Glu	CTT Leu	AGA Arg	ACA Thr 265	TTT Phe	GGG Gly	GGA Gly	CAT His	GAT Asp 270	GCA Ala	AAG Lys	816
TTT Phe	ATA Ile	GAT Asp 275	AGT Ser	TTA Leu	CAG Gln	GAA Glu	AAC Asn 280	GAA Glu	TTT Phe	CGT Arg	CTA	TAT Tyr 285	TAT Tyr	TAT Tyr	AAT Asn	864
						AGT Ser 295										912
GGT Gly 305	ACT Thr	ACT Thr	GCT Ala	TCA Ser	TTA Leu 310	CAG Gln	TAT Tyr	ATG Met	AAA Lys	AAT Asn 315	GTT Val	TTT Phe	AAA Lys	GAG Glu	AAA Lys 320	960
TAT Tyr	CTC Leu	CTA Leu	TCT Ser	GAA Glu 325	GAT Asp	ACA Thr	TCT Ser	GGA Gly	AAA Lys 330	TTT Phe	TCG Ser	GTA Val	GAT Asp	AAA Lys 335	TTA Leu	1008
						AAA Lys										1056
						AAA Lys										1104
						AAG Lys 375										1152
						AAT Asn										1200
TTT Phe	AAT Asn	GGT Gly	CAA Gln	AAT Asn 405	ACA Thr	GAA Glu	ATT Ile	AAT Asn	AAT Asn 410	ATG Met	AAT Asn	TTT Phe	ACT Thr	AAA Lys 415	CTA Leu	1248
						TTT Phe										1296
						ACT Thr										1344
						ATC Ile 455										1392

AGT Ser 465	Pro	TCA Ser	GAA Glu	GAT Asp	AAT Asn 470	TTT Phe	ACT Thr	AAT Asn	GAT Asp	CTA Leu 475	AAT Asn	AAA Lys	GGA Gly	GAA Glu	GAA Glu 480	1440
ATT Ile	ACA Thr	TCT Ser	GAT Asp	ACT Thr 485	AAT Asn	ATA Ile	GAA Glu	GCA Ala	GCA Ala 490	GAA Glu	GAA Glu	AAT Asn	ATT Ile	AGT Ser 495	TTA Leu	1488
						TAT Tyr										1536
GAA Glu	AAT Asn	ATT Ile 515	Ser	ATA Ile	GAA Glu	AAT Asn	CTT Leu 520	TCA Ser	AGT Ser	GAC Asp	ATT Ile	ATA Ile 525	GGC Gly	CAA Gln	TTA Leu	1584
GAA Glu	CTT Leu 530	ATG Met	CCT Pro	AAT Asn	ATA Ile	GAA Glu 535	AGA Arg	TTT Phe	CCT Pro	AAT Asn	GGA Gly 540	AAA Lys	AAG Lys	TAT Tyr	GAG Glu	1632
						TTC Phe										1680
						GCT Ala										1728
						TAT Tyr										1776
						GAG Glu										1824
						ACC Thr 615										1872
						ACT Thr										1920
						TTA Leu										1968
		Ser				ATT Ile										2016
						TTT Phe										2064
						ATA Ile 695										2112

AAA Lys 705	TGG Trp	GAT Asp	GAG Glu	GTC Val	TAT Tyr 710	AAA Lys	TAT Tyr	ATA Ile	GTA Val	ACA Thr 715	AAT Asn	TGG Trp	TTA Leu	GCA Ala	AAG Lys 720	2160
GTT Val	AAT Asn	ACA Thr	CAG Gln	ATT Ile 725	GAT Asp	CTA Leu	ATA Ile	AGA Arg	AAA Lys 730	AAA Lys	ATG Met	AAA Lys	GAA Glu	GCT Ala 735	TTA Leu	2208
GAA Glu	AAT Asn	CAA Gln	GCA Ala 740	GAA Glu	GCA Ala	ACA Thr	AAG Lys	GCT Ala 745	ATA Ile	ATA Ile	AAC Asn	TAT Tyr	CAG Gln 750	TAT Tyr	AAT Asn	2256
						AAA Lys										2304
						GAG Glu 775										2352
						TGC Cys										2400
						CGG Arg										2448
						ATA Ile										2496
						GAT Asp										2544
						AAA Lys 855										2592
						AAG Lys										2640
						CAT His										2688
						AAA Lys										2736
						TTA Leu										2784
						AAT Asn 935										2832

TTT TGG ATA Phe Trp Ile 945	AGA ATT CCT Arg Ile Pro 950	Lys Tyr Phe	r AAC AGT ATA Asn Ser Ile 955	A AGT CTA AAT e Ser Leu Asn	AAT 2880 Asn 960
GAA TAT ACA Glu Tyr Thr	ATA ATA AAT Ile Ile Asn 965	TGT ATG GAP Cys Met Glu	A AAT AAT TCA 1 Asn Asn Sei 970	A GGA TGG AAA r Gly Trp Lys 975	GTA 2928 Val
TCA CTT AAT Ser Leu Asn	TAT GGT GAA Tyr Gly Glu 980	ATA ATC TGG Ile Ile Trp 985	Thr Leu Glr	G GAT ACT CAG n Asp Thr Gln 990	GAA 2976 Glu
ATA AAA CAA Ile Lys Gln 995	Arg Val Val	TTT AAA TAC Phe Lys Tyr 1000	C AGT CAA ATO C Ser Gln Met	G ATT AAT ATA t Ile Asn Ile 1005	TCA 3024 Ser
GAT TAT ATA Asp Tyr Ile 1010	AAC AGA TGG Asn Arg Trp	ATT TTT GTA Ile Phe Val 1015	A ACT ATC ACT Thr Ile Thr 102	r AAT AAT AGA r Asn Asn Arg 20	TTA 3072 Leu
AAT AAC TCT Asn Asn Ser 1025	AAA ATT TAT Lys Ile Tyr 103	Ile Asn Gly	A AGA TTA ATA Arg Leu Ile 1035	A GAT CAA AAA e Asp Gln Lys	CCA 3120 Pro 1040
ATT TCA AAT Ile Ser Asn	TTA GGT AAT Leu Gly Asn 1045	ATT CAT GCT Ile His Ala	AGT AAT AAT Ser Asn Asr 1050	r ATA ATG TTT n Ile Met Phe 105	Lys
TTA GAT GGT Leu Asp Gly	TGT AGA GAT Cys Arg Asp 1060	ACA CAT AGA Thr His Arg 106	Tyr Ile Tr	G ATA AAA TAT O Ile Lys Tyr 1070	TTT 3216 Phe
AAT CTT TTT Asn Leu Phe 107	Asp Lys Glu	TTA AAT GAA Leu Asn Glu 1080	AAA GAA ATO Lys Glu Ile	C AAA GAT TTA E Lys Asp Leu 1085	TAT 3264 Tyr
GAT AAT CAA Asp Asn Gln 1090	TCA AAT TCA Ser Asn Ser	GGT ATT TTA Gly Ile Leu 1095	A AAA GAC TTT Lys Asp Phe 110	T TGG GGT GAT Trp Gly Asp	TAT 3312 Tyr
TTA CAA TAT Leu Gln Tyr 1105	GAT AAA CCA Asp Lys Pro 1110	Tyr Tyr Met	TTA AAT TTA Leu Asn Leu 1115	A TAT GAT CCA 1 Tyr Asp Pro	AAT 3360 Asn 1120
AAA TAT GTC Lys Tyr Val	GAT GTA AAT Asp Val Asn 1125	AAT GTA GGT Asn Val Gly	TATT AGA GGT Ile Arg Gly 1130	T TAT ATG TAT Y Tyr Met Tyr 113	Leu
AAA GGG CCT Lys Gly Pro	AGA GGT AGC Arg Gly Ser 1140	GTA ATG ACT Val Met Thr 114	Thr Asn Ile	T TAT TTA AAT E Tyr Leu Asn 1150	TCA 3456 Ser
AGT TTG TAT Ser Leu Tyr 115	Arg Gly Thr	AAA TTT ATT Lys Phe Ile 1160	TATA AAA AAA E Ile Lys Lys	A TAT GCT TCT s Tyr Ala Ser 1165	GGA 3504 Gly
AAT AAA GAT Asn Lys Asp 1170	AAT ATT GTT Asn Ile Val	AGA AAT AAT Arg Asn Asn 1175	GAT CGT GTA Asp Arg Val	A TAT ATT AAT l Tyr Ile Asn 80	GTA 3552 Val

	GTA Val 118!	Val	AAA Lys	AAT Asn	AAA Lys	GAA Glu 119	Tyr	AGG Arg	TTA Leu	GCT Ala	ACT Thr 119	Asn	GCA Ala	TCA Ser	CAG Gln	GCA Ala 1200	36	500
	GGC Gly	GTA Val	GAA Glu	AAA Lys	ATA Ile 120	Leu	AGT Ser	GCA Ala	TTA Leu	GAA Glu 121	Ile	CCT Pro	GAT Asp	GTA Val	GGA Gly 121	Asn	36	548
:	CTA Leu	AGT Ser	CAA Gln	GTA Val 1220	Val	GTA Val	ATG Met	AAG Lys	TCA Ser 1225	Lys	AAT Asn	GAT Asp	CAA Gln	GGA Gly 1230	Ile	ACA Thr	36	96
1	AAT Asn	AAA Lys	TGC Cys 1235	Lys	ATG Met	AAT Asn	TTA Leu	CAA Gln 1240	Asp	AAT Asn	AAT Asn	GGG Gly	AAT Asn 1245	Asp	ATA Ile	GGC Gly	37	44
]	rrr Phe	ATA Ile 1250	Gly	TTT Phe	CAT His	CAG Glņ	TTT Phe 1255	Asn	AAT Asn	ATA Ile	GCT Ala	AAA Lys 1260	Leu	GTA Val	GCA Ala	AGT Ser	37	92
1	AAT Asn L265	Trp	TAT Tyr	AAT Asn	AGA Arg	CAA Gln 1270	ATA Ile	GAA Glu	AGA Arg	TCT Ser	AGT Ser 1275	Arg	ACT Thr	TTG Leu	GGT Gly	TGC Cys 1280	38	40
7	CA Ser	TGG Trp	GAA Glu	Phe	ATT Ile 1285	Pro	GTA Val	GAT Asp	Asp	GGA Gly 1290	Trp'	GGA Gly	GAA Glu	AGG Arg	CCA Pro 1295	Leu	38	88
2	raa																38	91
•	(2)	INFO	RMAT	NOI	FOR	SEQ	ID N	0:28	:									

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1296 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:28:

Met Gln Phe Val Asn Lys Gln Phe Asn Tyr Lys Asp Pro Val Asn Gly
1 5 10 15

Val Asp Ile Ala Tyr Ile Lys Ile Pro Asn Val Gly Gln Met Gln Pro 20 25 30

Val Lys Ala Phe Lys Ile His Asn Lys Ile Trp Val Ile Pro Glu Arg
35 40 45

Asp Thr Phe Thr Asn Pro Glu Glu Gly Asp Leu Asn Pro Pro Pro Glu 50 55 60

Ala Lys Gln Val Pro Val Ser Tyr Tyr Asp Ser Thr Tyr Leu Ser Thr 65 70 75 80

Asp Asn Glu Lys Asp Asn Tyr Leu Lys Gly Val Thr Lys Leu Phe Glu 85 90 95

Arg Ile Tyr Ser Thr Asp Leu Gly Arg Met Leu Leu Thr Ser Ile Val

Arg Gly Ile Pro Phe Trp Gly Gly Ser Thr Ile Asp Thr Glu Leu Lys Val Ile Asp Thr Asn Cys Ile Asn Val Ile Gln Pro Asp Gly Ser Tyr 135 Arg Ser Glu Glu Leu Asn Leu Val Ile Ile Gly Pro Ser Ala Asp Ile Ile Gln Phe Glu Cys Lys Ser Phe Gly His Glu Val Leu Asn Leu Thr 170 Arg Asn Gly Tyr Gly Ser Thr Gln Tyr Ile Arg Phe Ser Pro Asp Phe 185 Thr Phe Gly Phe Glu Glu Ser Leu Glu Val Asp Thr Asn Pro Leu Leu 200 Gly Ala Gly Lys Phe Ala Thr Asp Pro Ala Val Thr Leu Ala His Glu Leu Ile His Ala Gly His Arg Leu Tyr Gly Ile Ala Ile Asn Pro Asn Arg Val Phe Lys Val Asn Thr Asn Ala Tyr Tyr Glu Met Ser Gly Leu Glu Val Ser Phe Glu Glu Leu Arg Thr Phe Gly Gly His Asp Ala Lys Phe Ile Asp Ser Leu Gln Glu Asn Glu Phe Arg Leu Tyr Tyr Asn 280 Lys Phe Lys Asp Ile Ala Ser Thr Leu Asn Lys Ala Lys Ser Ile Val-Gly Thr Thr Ala Ser Leu Gln Tyr Met Lys Asn Val Phe Lys Glu Lys Tyr Leu Leu Ser Glu Asp Thr Ser Gly Lys Phe Ser Val Asp Lys Leu 325 Lys Phe Asp Lys Leu Tyr Lys Met Leu Thr Glu Ile Tyr Thr Glu Asp Asn Phe Val Lys Phe Phe Lys Val Leu Asn Arg Lys Thr Tyr Leu Asn Phe Asp Lys Ala Val Phe Lys Ile Asn Ile Val Pro Lys Val Asn Tyr 375 380 Thr Ile Tyr Asp Gly Phe Asn Leu Arg Asn Thr Asn Leu Ala Ala Asn Phe Asn Gly Gln Asn Thr Glu Ile Asn Asn Met Asn Phe Thr Lys Leu 405 410 Lys Asn Phe Thr Gly Leu Phe Glu Phe Tyr Lys Leu Leu Cys Val Arg 425 Gly Ile Ile Thr Ser Lys Thr Lys Ser Leu Asp Lys Gly Tyr Asn Lys

Ala Leu Asn Asp Leu Cys Ile Lys Val Asn Asn Trp Asp Leu Phe Phe Ser Pro Ser Glu Asp Asn Phe Thr Asn Asp Leu Asn Lys Gly Glu Glu 470 Ile Thr Ser Asp Thr Asn Ile Glu Ala Ala Glu Glu Asn Ile Ser Leu Asp Leu Ile Gln Gln Tyr Tyr Leu Thr Phe Asn Phe Asp Asn Glu Pro Glu Asn Ile Ser Ile Glu Asn Leu Ser Ser Asp Ile Ile Gly Gln Leu 520 Glu Leu Met Pro Asn Ile Glu Arg Phe Pro Asn Gly Lys Lys Tyr Glu Leu Asp Lys Tyr Thr Met Phe His Tyr Leu Arg Ala Gln Glu Phe Glu 555 His Gly Lys Ser Arg Ile Ala Leu Thr Asn Ser Val Asn Glu Ala Leu 570 Leu Asn Pro Ser Arg Val Tyr Thr Phe Phe Ser Ser Asp Tyr Val Lys Lys Val Asn Lys Ala Thr Glu Ala Ala Met Phe Leu Gly Trp Val Glu 600 Gln Leu Val Tyr Asp Phe Thr Asp Glu Thr Ser Glu Val Ser Thr Thr Asp Lys Ile Ala Asp Ile Thr Ile Ile Ile Pro Tyr Ile Gly Pro Ala 630 Leu Asn Ile Gly Asn Met Leu Tyr Lys Asp Asp Phe Val Gly Ala Leu Ile Phe Ser Gly Ala Val Ile Leu Leu Glu Phe Ile Pro Glu Ile Ala Ile Pro Val Leu Gly Thr Phe Ala Leu Val Ser Tyr Ile Ala Asn Lys 680 Val Leu Thr Val Gln Thr Ile Asp Asn Ala Leu Ser Lys Arg Asn Glu 695 Lys Trp Asp Glu Val Tyr Lys Tyr Ile Val Thr Asn Trp Leu Ala Lys 715 Val Asn Thr Gln Ile Asp Leu Ile Arg Lys Lys Met Lys Glu Ala Leu Glu Asn Gln Ala Glu Ala Thr Lys Ala Ile Ile Asn Tyr Gln Tyr Asn Gln Tyr Thr Glu Glu Glu Lys Asn Asn Ile Asn Phe Asn Ile Asp Asp 760 Leu Ser Ser Lys Leu Asn Glu Ser Ile Asn Lys Ala Met Ile Asn Ile

Asn Lys Phe Leu Asn Gln Cys Ser Val Ser Tyr Leu Met Asn Ser Met Ile Pro Tyr Gly Val Lys Arg Leu Glu Asp Phe Asp Ala Ser Leu Lys Asp Ala Leu Leu Lys Tyr Ile Tyr Asp Asn Arg Gly Thr Leu Ile Gly Gln Val Asp Arg Leu Lys Asp Lys Val Asn Asn Thr Leu Ser Thr Asp 840 Ile Pro Phe Gln Leu Ser Lys Tyr Val Asp Asn Gln Arg Leu Leu Ser 850 Thr Phe Thr Glu Tyr Ile Lys Asn Ile Ile Asn Thr Ser Ile Leu Asn 875 870 Leu Arg Tyr Glu Ser Asn His Leu Ile Asp Leu Ser Arg Tyr Ala Ser Lys Ile Asn Ile Gly Ser Lys Val Asn Phe Asp Pro Ile Asp Lys Asn 905 Gln Ile Gln Leu Phe Asn Leu Glu Ser Ser Lys Ile Glu Val Ile Leu Lys Asn Ala Ile Val Tyr Asn Ser Met Tyr Glu Asn Phe Ser Thr Ser Phe Trp Ile Arg Ile Pro Lys Tyr Phe Asn Ser Ile Ser Leu Asn Asn Glu Tyr Thr Ile Ile Asn Cys Met Glu Asn Asn Ser Gly Trp Lys Val Ser Leu Asn Tyr Gly Glu Ile Ile Trp Thr Leu Gln Asp Thr Gln Glu Ile Lys Gln Arg Val Val Phe Lys Tyr Ser Gln Met Ile Asn Ile Ser 1000 995 Asp Tyr Ile Asn Arg Trp Ile Phe Val Thr Ile Thr Asn Asn Arg Leu 1020 Asn Asn Ser Lys Ile Tyr Ile Asn Gly Arg Leu Ile Asp Gln Lys Pro 1035 Ile Ser Asn Leu Gly Asn Ile His Ala Ser Asn Asn Ile Met Phe Lys 1045 Leu Asp Gly Cys Arg Asp Thr His Arg Tyr Ile Trp Ile Lys Tyr Phe 1065 Asn Leu Phe Asp Lys Glu Leu Asn Glu Lys Glu Ile Lys Asp Leu Tyr 1080 1075

Asp Asn Gln Ser Asn Ser Gly Ile Leu Lys Asp Phe Trp Gly Asp Tyr

Leu Gln Tyr Asp Lys Pro Tyr Tyr Met Leu Asn Leu Tyr Asp Pro Asn

1095

1110

1105

1115

1100

		•													
Lys	Tyr	Val	Asp	Val 112		Asn	Val	Gly	Ile 113		Gly	Tyr	Met	Tyr 113	
Lys	Gly	Pro	Arg		Ser	Val	Met	Thr 114		Asn	Ile	Tyr	Leu 115		Ser
Ser	Leu	Tyr 115		Gly	Thr	Lys	Phe 116		Ile	Lys	Lys	Tyr 116		Ser	Gly
Asn	Lys 117		Asn	Ile	Val	Arg 1179		Asn	Asp	Arg	Val 1180		Ile	Asn	Val
Val 1185	Val	Lys	Asn	Lys	Glu 1190	Tyr O	Arg	Leu	Ala	Thr 1199		Ala	Ser	Gln	Ala 1200
Gly	Val	Glu	Lys	Ile 1205		Ser	Ala	Leu	Glu 121		Pro	Asp	Val	Gly 121	
Leu	Ser	Gln	Val 1220		Val	Met	Lys	Ser 1225		Asn	Asp	Gln	Gly 1230		Thr
Asn	Lys	Cys 1235		Met	Asn	Leu	Gln 1240		Asn	Asn	Gly	Asn 1245	_	Ile	Gly
Phe	Ile 1250	-	Phe	His	Gln	Phe 1255		Asn	Ile	Ala	Lys 1260		Val	Ala	Ser
Asn 1265		Tyr	Asn	Arg	Gln 1270	Ile	Glu	Arg	Ser	Ser 1275	_	Thr	Leu	Gly	Cys 1280
Ser	Trp	Glu	Phe	Ile 1285		Val	Asp	Asp	Gly 1290	-	Gly	Glu	Arg	Pro 1299	
(2)	INFO	RMAT	NOI	FOR	SEQ	ID N	10:29):		,					
	(i)	(A (B (C	L) LE () TY () ST	NGTH PE: RAND	: 30 nucl EDNE	TERI bas eic SS: line	e pa acid sing	irs l							
	(ii)					othe									
	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	:29:					
CGCC	ATGG	CT A	GATT	'ATTA	т ст	'ACAT	TTAC								
(2)	INFO	RMAT	ION	FOR	SEQ	ID N	0:30	:							
	(i)					TERI									

- (A) LENGTH: 26 base pairs(B) TYPE: nucleic acid(C) STRANDEDNESS: single(D) TOPOLOGY: linear

- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:30:

GCAAGCTTCT TGACAGACTC ATGTAG

(2) INFORMATION FOR SEQ ID NO:31:

- (i) SEQUENCE CHARACTERISTICS:
 (A) LENGTH: 1546 base pairs
 (B) TYPE: nucleic acid
 (C) STRANDEDNESS: double

 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:31:

AGATCTCGAT	CCCGCGAAAT	TAATACGACT	CACTATAGGG	GAATTGTGAG	CGGATAACAA	60
TTCCCCTCTA	GAAATAATTT	TGTTTAACTT	TAAGAAGGAG	ATATACCATG	GGCCATCATC	120
ATCATCATCA	TCATCATCAT	CACAGCAGCG	GCCATATCGA	AGGTCGTCAT	ATGGCTAGCA	180
TGGCTAGATT	ATTATCTACA	TTTACTGAAT	ATATTAAGAA	TATTATTAAT	ACTTCTATAT	240
TGAATTTAAG	ATATGAAAGT	AATCATTTAA	TAGACTTATC	TAGGTATGCA	тсааааатаа	300
ATATTGGTAG	TAAAGTAAAT	TTTGATCCAA	TAGATAAAAA	TCAAATTCAA	TTATTTAATT	360
TAGAAAGTAG	TAAAATTGAG	GTAATTTTAA	AAAATGCTAT	TGTATATAAT	AGTATGTATG	420
AAAATTTTAG	TACTAGCTTT	TGGATAAGAA	TTCCTAAGTA	TTTTAACAGT	ATAAGTCTAA	480
ATAATGAATA	ТАСААТААТА	AATTGTATGG	AAAATAATTC	AGGATGGAAA	GTATCACTTA	540
ATTATGGTGA	AATAATCTGG	ACTTTACAGG	ATACTCAGGA	AATAAAACAA	AGAGTAGTTT	600
TTAAATACAG	TCAAATGATT	AATATATCAG	ATTATATAAA	CAGATGGATT	TTTGTAACTA	660
TCACTAATAA	TAGATTAAAT	ААСТСТАААА	TTTATATAAA	TGGAAGATTA	ATAGATCAAA	720
AACCAATTTC	AAATTTAGGT	AATATTCATG	CTAGTAATAA	TATAATGTTT	AAATTAGATG	780
GTTGTAGAGA	TACACATAGA	TATATTTGGA	TAAAATATTT	TAATCTTTTT	GATAAGGAAT	840
TAAATGAAAA	AGAAATCAAA	GATTTATATG	ATAATCAATC	AAATTCAGGT	ATTTTAAAAG	900
ACTTTTGGGG	TGATTATTTA	CAATATGATA	AACCATACTA	TATGTTAAAT	TTATATGATC	960
CAAATAAATA	TGTCGATGTA	AATAATGTAG	GTATTAGAGG	TTATATGTAT	CTTAAAGGGC	1020
CTAGAGGTAG	CGTAATGACT	ACAAACATTT	ATTTAAATTC	AAGTTTGTAT	AGGGGGACAA	1080
AATTTATTAT	ААААААТАТ	GCTTCTGGAA	ATAAAGATAA	TATTGTTAGA	AATAATGATC	1140
GTGTATATAT	TAATGTAGTA	GTTAAAAATA	AAGAATATAG	GTTAGCTACT	AATGCATCAC	1200
AGGCAGGCGT	AGAAAAAATA	CTAAGTGCAT	TAGAAATACC	TGATGTAGGA	AATCTAAGTC	1260
AAGTAGTAGT	AATGAAGTCA	AAAAATGATC	AAGGAATAAC	AAATAAATGC	AAAATGAATT	1320
TACAAGATAA	TAATGGGAAT	GATATAGGCT	TTATAGGATT	TCATCAGTTT	AATAATATAG	1380
CTAAACTAGT	AGCAAGTAAT	TGGTATAATA	GACAAATAGA	AAGATCTAGT	AGGACTTTGG	1440

GT	GUTCATG GGAATITATT CCTGTAGATG ATGGATGGGG AGAAAGGCCA CTGTAATTAA	1500
TCT	CAAACTA CATGAGTCTG TCAAGAAGCT TGCGGCCGCA CTCGAG	1546
(2)	INFORMATION FOR SEQ ID NO:32:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 9 amino acids (B) TYPE: amino acid (C) STRANDEDNESS: not relevant (D) TOPOLOGY: not relevant 	
	(ii) MOLECULE TYPE: peptide	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:32:	
	Met His His His His Met Ala 1 5	
(2)	INFORMATION FOR SEQ ID NO:33:	
	 (i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 21 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear 	
	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "DNA"</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:33:	
TAT	GCATCAC CATCACCATC A	21
(2)	INFORMATION FOR SEQ ID NO:34:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 23 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single (D) TOPOLOGY: linear	
	<pre>(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "DNA"</pre>	
	(xi) SEQUENCE DESCRIPTION: SEQ ID NO:34:	
CAT	ETGATGG TGATGGTGAT GCA	23
(2)	INFORMATION FOR SEQ ID NO:35:	
	(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 1351 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: double (D) TOPOLOGY: linear	
	(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "DNA"	٠.

(ix) FEATURE:
(A) NAME/KEY: CDS
(B) LOCATION: 1..1335

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:35:

	•		2000				011.	224		0.33	•						
ATG Met 1	His	CAC His	CAT His	CAC His 5	CAT His	CAC His	ATG Met	GCT Ala	CGT Arg 10	Leu	CTG Leu	TCT Ser	ACC Thr	TTC Phe 15	ACT Thr		48
GAA Glu	TAC	ATC	AAG Lys 20	AAC Asn	ATC Ile	ATC Ile	AAT Asn	ACC Thr 25	TCC Ser	ATC Ile	CTG Leu	AAC Asn	CTG Leu 30	CGC Arg	TAC Tyr	•	96
GAA Glu	TCC Ser	AAT Asn 35	CAC His	CTG Leu	ATC Ile	GAC Asp	CTG Leu 40	TCT Ser	CGC Arg	TAC Tyr	Ala	TCC Ser 45	AAA Lys	ATC Ile	AAC Asn		144
ATC Ile	GGT Gly 50	TCT	AAA Lys	GTT Val	AAC Asn	TTC Phe 55	GAT Asp	CCG Pro	ATC Ile	GAC Asp	AAG Lys 60	AAT Asn	CAG Gln	ATC Ile	CAG Gln		192
CTG Leu 65	TTC Phe	AAT Asn	CTG Leu	GAA Glu	TCT Ser 70	TCC Ser	AAA Lys	ATC Ile	GAA Glu	GTT Val 75	ATC Ile	CTG Leu	AAG Lys	AAT Asn	GCT Ala 80		240
ATC Ile	GTA Val	TAC Tyr	AAC Asn	TCT Ser 85	ATG Met	TAC Tyr	GAA Glu	AAC Asn	TTC Phe 90	TCC Ser	ACC Thr	TCC Ser	TTC Phe	TGG Trp 95	ATC Ile		288
CGT Arg	ATC Ile	CCG Pro	AAA Lys 100	TAC Tyr	TTC Phe	AAC Asn	TCC Ser	ATC Ile 105	TCT Ser	CTG Leu	AAC Asn	AAT Asn	GAA Glu 110	TAC Tyr	ACC Thr		336
		AAC Asn 115									Lys						384
		GAA Glu															432
		GTA Val															480
		TGG Trp															528
		TAC Tyr														•	576
		AAC Asn 195															624
TGT Cys		GAC Asp															672

GAC Asp 225	Lys	GAA Glu	CTG Leu	AAC Asn	GAA Glu 230	AAA Lys	GAA Glu	ATC Ile	AAA Lys	GAC Asp 235	CTG Leu	TAC Tyr	GAC Asp	AAC Asn	CAG Gln 240	72	0
TCC Ser	AAT Asn	TCT Ser	GGT Gly	ATC Ile 245	CTG Leu	AAA Lys	GAC Asp	TTC Phe	TGG Trp 250	GGT Gly	GAC Asp	TAC Tyr	CTG Leu	CAG Gln 255	TAC Tyr	76	8
									TAC Tyr							81	6
									TAC Tyr							86	4
									TAC Tyr							91:	2
CGT Arg 305	GGT Gly	ACC Thr	AAA Lys	TTC Phe	ATC Ile 310	ATC Ile	AAG Lys	AAA Lys	TAC Tyr	GCG Ala 315	TCT Ser	GGT Gly	AAC Asn	AAG Lys	GAC Asp 320	960	5
									TAC Tyr 330							1008	3
									GCT Ala							1056	5
									GAC Asp							1104	1
									CAG Gln							1152	2
				Gln					AAC Asn							1200)
TTC Phe	CAC His	CAG Gln	TTC Phe	AAC Asn 405	AAT Asn	ATC Ile	GCT Ala	AAA Lys	CTG Leu 410	GTT Val	GCT Ala	TCC Ser	AAC Asn	TGG Trp 415	TAC Tyr	1248	3
									ACT Thr							1296	5
									GAA Glu				TAAC	CCGG	GA .	1345	5
AAGO	TT															1351	L

(2) INFORMATION FOR SEQ ID NO:36:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 445 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:36:

Met His His His His His Met Ala Arg Leu Leu Ser Thr Phe Thr
1 10 15

Glu Tyr Ile Lys Asn Ile Ile Asn Thr Ser Ile Leu Asn Leu Arg Tyr
20 25 30

Glu Ser Asn His Leu Ile Asp Leu Ser Arg Tyr Ala Ser Lys Ile Asn 35 40 45

Ile Gly Ser Lys Val Asn Phe Asp Pro Ile Asp Lys Asn Gln Ile Gln
50 55 60

Leu Phe Asn Leu Glu Ser Ser Lys Ile Glu Val Ile Leu Lys Asn Ala 65 70 75 80

Ile Val Tyr Asn Ser Met Tyr Glu Asn Phe Ser Thr Ser Phe Trp Ile
85 90 95

Arg Ile Pro Lys Tyr Phe Asn Ser Ile Ser Leu Asn Asn Glu Tyr Thr
100 105 110

Ile Ile Asn Cys Met Glu Asn Asn Ser Gly Trp Lys Val Ser Leu Asn 115 120 125

Tyr Gly Glu Ile Ile Trp Thr Leu Gln Asp Thr Gln Glu Ile Lys Gln 130 135 140

Arg Val Val Phe Lys Tyr Ser Gln Met Ile Asn Ile Ser Asp Tyr Ile 145 150 155 160

Asn Arg Trp Ile Phe Val Thr Ile Thr Asn Asn Arg Leu Asn Asn Ser 165 170 175

Lys Ile Tyr Ile Asn Gly Arg Leu Ile Asp Gln Lys Pro Ile Ser Asn 180 185 190

Leu Gly Asn Ile His Ala Ser Asn Asn Ile Met Phe Lys Leu Asp Gly
195 200 205

Cys Arg Asp Thr His Arg Tyr Ile Trp Ile Lys Tyr Phe Asn Leu Phe 210 215 220

Asp Lys Glu Leu Asn Glu Lys Glu Ile Lys Asp Leu Tyr Asp Asn Gln 225 230 235 240

Ser Asn Ser Gly Ile Leu Lys Asp Phe Trp Gly Asp Tyr Leu Gln Tyr 245 250 255

Asp Lys Pro Tyr Tyr Met Leu Asn Leu Tyr Asp Pro Asn Lys Tyr Val 260 265 270

Asp	Val	Asn 275	Asn	Val	Gly	Ile	Arg 280	Gly	Tyr	Met	Tyr	Leu 285	Lys	Gly	Pro		
Arg	Gly 290	Ser	Val	Met	Thr	Thr 295	Asn	Ile	Tyr	Leu	Asn 300	Ser	Ser	Leu	Tyr		
Arg 305		Thr	Lys	Phe	Ile 310	Ile	Lys	Lys	Tyr	Ala 315	Ser	Gly	Asn	Lys	Asp 320		
Asn	Ile	Val	Arg	Asn 325	Asn	Asp	Arg	Val	Tyr 330	Ile	Asn	Val	Val	Val 335	Lys		
Asn	Lys	Glu	Tyr 340	Arg	Leu	Ala	Thr	Asn 345	Ala	Ser	Gln	Ala	Gly 350	Val	Glu		
Lys	Ile	Leu 355	Ser	Ala	Leu	Glu	Ile 360	Pro	Asp	Val	Gly	Asn 365	Leu	Ser	Gln		
Val	Val 370	Val	Met	Lys	Ser	Lys 375	Asn	Asp	Gln	Gly	Ile 380	Thr	Asn	Lys	Cys		
Lys 385	Met	Asn	Leu	Gln	Asp 390	Asn	Asn	Gly	Asn	Asp 395	Ile	Gly	Phe	Ile	Gly 400	•	
Phe	His	Gln	Phe	Asn 405	Asn	Ile	Ala	Lys	Leu 410	Val	Ala	Ser	Asn	Trp 415	Tyr		
Asn	Arg	Gln	Ile 420	Glu	Arg	Ser	Ser	Arg 425	Thr	Leu	Gly	Cys	Ser 430	Trp	Glu		
Phe	Ile	Pro 435	Val	Asp	Asp	Gly	Trp 440	Gly	Glu	Arg	Pro	Leu 445					
(2)	INFO	ORMAT	CION	FOR	SEQ	ID N	10:37	7:									
	(i) (ii)	(A) (E) (C) (D) MOL	A) LE B) TY C) ST D) TO LECUI	ENGTH (PE: TRANI DPOLO LE TY	I: 27 nucl EDNE GY:		e pa acid sing ar	airs l gle									
	(xi)	SEÇ			•												
CGC	ATATO	SAA I	ATTO	GTC	A TI	GCAT	'G										27
(2)	INFO	RMAT	CION	FOR	SEQ	ID N	O:38	3:									
		. (E	L) LE B) TY C) ST	ENGTI (PE : TRANI	I: 27 nucl EDNE	bas eic	e pa acid sing	irs i									
	(ii)	MOI (P	ECUI	E TY	PE:	othe N:/	r nu desc	clei = "	c ac	id'							
	(xi)	SEC	UENC	E DE	SCRI	PTIC	N: S	SEQ I	D NC	:38							

27

GGAAGCTTGC AGGGCAATTA CATCATG

	(:		(A) 1 (B) 1 (C) 5	LENG? FYPE : STRAN	TH: 3 : nuc VDEDN	3876 Cleid	base aci dou	e pa: id	irs							
	(ii	L) MC	LECT	TE 1	YPE:	DNA	(ge	enomi	ic)							
	(i)		(A)	IAME/		CDS	; 3873	}								
		.) SE														
ATG Met	Pro	GTI Val	ACA Thr	ATA Ile 5	Asn	AAT Asn	TTT Phe	AAT Asn	TAT Tyr 10	Asn	GAT Asp	Pro	ATT Ile	GAT Asp 15	AAT Asn	4.8
GAC Asp	AAT Asn	ATT	ATT Ile 20	Met	ATG Met	GAA Glu	CCT Pro	CCA Pro 25	Phe	GCA Ala	AGG Arg	GGT Gly	ACG Thr	Gly	AGA Arg	96
TAT Tyr	TAT Tyr	AAA Lys 35	Ala	TTT Phe	AAA Lys	ATC Ile	ACA Thr 40	GAT Asp	CGT	ATT	TGG Trp	ATA Ile 45	ATA Ile	CCC	GAA Glu	144
AGA Arg	TAT Tyr 50	ACT Thr	TTT Phe	GGA Gly	TAT Tyr	AAA Lys 55	CCT	GAG Glu	GAT Asp	TTT Phe	AAT Asn 60	AAA Lys	AGT Ser	TCC	GGT Gly	192
ATT Ile 65	TTT Phe	AAT Asn	AGA Arg	GAT Asp	GTT Val 70	TGT Cys	GAA Glu	TAT Tyr	TAT Tyr	GAT Asp 75	CCA Pro	GAT Asp	TAC Tyr	TTA Leu	AAT Asn 80	240
ACC Thr	AAT Asn	GAT Asp	AAA Lys	AAG Lys 85	AAT Asn	ATA Ile	TTT Phe	TTC Phe	CAA Gln 90	ACA Thr	TTG Leu	ATC Ile	AAG Lys	TTA Leu 95	TTT Phe	288
AAT Asn	AGA Arg	ATC Ile	AAA Lys 100	TCA Ser	AAA Lys	CCA Pro	TTG Leu	GGT Gly 105	GAA Glu	AAG Lys	TTA Leu	TTA Leu	GAG Glu 110	ATG Met	ATT Ile	336
ATA Ile	Asn	GGT Gly 115	ATA Ile	CCT Pro	TAT Tyr	CTT Leu	GGA Gly 120	GAT Asp	AGA Arg	CGT Arg	GTT Val	CCA Pro 125	CTC Leu	GAA Glu	GAG Glu	384
TTT Phe	AAC Asn 130	ACA Thr	AAC Asn	ATT Ile	GCT Ala	AGT Ser 135	GTA Val	ACT Thr	GTT Val	AAT Asn	AAA Lys 140	TTA Leu	ATT Ile	AGT Ser	AAT Asn	432
CCA Pro 145	GGA Gly	GAA Glu	GTG Val	GAG Glu	CGA Arg 150	AAA Lys	aaa Lys	GGT Gly	ATT Ile	TTC Phe 155	GCA Ala	AAT Asn	TTA Leu	ATA Ile	ATA Ile 160	480
TTT Phe	GGA Gly	CCT Pro	GGG Gly	CCA Pro 165	GTT Val	TTA Leu	AAT Asn	GAA Glu	AAT Asn 170	GAG Glu	ACT Thr	ATA Ile	GAT Asp	ATA Ile 175	GGT Gly	528

(2) INFORMATION FOR SEQ ID NO:39:

ATA Ile	CAA Gln	AAT Asn	CAT His 180	Phe	GCA Ala	TCA Ser	AGG Arg	GAA Glu 185	Gly	TTT	GGG Gly	GGT Gly	ATA Ile 190	Met	CAA Gln	576
ATG Met	AAA Lys	TTT Phe 195	Cys	CCA Pro	GAA Glu	TAT	GTA Val 200	Ser	GTA Val	TTT	AAT Asn	AAT Asn 205	GTT Val	CAA Gln	GAA Glu	624
AAC Asn	AAA Lys 210	Gly	GCA Ala	AGT Ser	ATA Ile	TTT Phe 215	AAT Asn	AGA Arg	CGT	GGA Gly	TAT Tyr 220	TTT Phe	TCA Ser	GAT Asp	CCA Pro	672
GCC Ala 225	TTG Leu	ATA Ile	TTA Leu	ATG Met	CAT His 230	Glu	CTT Leu	ATA Ile	CAT His	GTT Val 235	TTG Leu	CAT His	GGA Gly	TTA Leu	TAT Tyr 240	720
GGC Gly	ATT Ile	AAA Lys	GTA Val	GAT Asp 245	Asp	TTA Leu	CCA Pro	ATT Ile	GTA Val 250	CCA Pro	AAT Asn	GAA Glu	AAA Lys	AAA Lys 255	TTT Phe	768
TTT Phe	ATG Met	CAA Gln	TCT Ser 260	ACA Thr	GAT Asp	ACT	ATA Ile	CAG Gln 265	GCA Ala	GAA Glu	GAA Glu	CTA Leu	TAT Tyr 270	ACA Thr	TTT Phe	816
GGA Gly	GGA Gly	CAA Gln 275	GAT Asp	CCC	AGC Ser	ATC Ile	ATA Ile 280	TCT Ser	CCT Pro	TCT Ser	ACA Thr	GAT Asp 285	AAA Lys	AGT Ser	ATC Ile	864
						AAT Asn 295										912
AAG Lys 305	GTT Val	TTA Leu	GTT Val	TGC Cys	ATA Ile 310	TCA Ser	GAT Asp	CCT Pro	AAC Asn	ATT Ile 315	AAC Asn	ATT Ile	AAT Asn	ATA Ile	TAT Tyr 320	960
						AAA Lys										1008
						GAA Glu										1056
						ATT Ile										1104
		Ala				AGT Ser 375										1152
						ATC Ile										1200
						AAA Lys										1248

AAT Asn	AAA Lys	CAA Gln	GCT Ala 420	Tyr	GAA Glu	GAA Glu	ATC	AGC Ser 425	Lys	GAG Glu	CAT His	TTG Leu	GCT Ala 430	GTA Val	TAT	1296
AAG Lys	ATA Ile	CAA Gln 435	Met	TGT Cys	AAA Lys	AGT Ser	GTT Val 440	Lys	GTT Val	CCA Pro	GGA Gly	ATA Ile 445	TGT Cys	ATT	GAT Asp	1344
GTC Val	GAT Asp 450	Asn	GAA Glu	AAT Asn	TTG Leu	TTC Phe 455	TTT Phe	ATA Ile	GCT Ala	GAT Asp	AAA Lys 460	Asn	AGT Ser	TTT Phe	TCA Ser	1392
GAT Asp 465	Asp	TTA Leu	TCT Ser	AAA Lys	AAT Asn 470	GAA Glu	AGA Arg	GTA Val	GAA Glu	TAT Tyr 475	AAT Asn	ACA Thr	CAG Gln	AAT Asn	AAT Asn 480	1440
TAT Tyr	ATA Ile	GGA Gly	AAT Asn	GAC Asp 485	TTT	CCT	ATA Ile	AAT Asn	GAA Glu 490	TTA Leu	ATT Ile	TTA Leu	GAT Asp	ACT Thr 495	GAT Asp	1488
TTA Leu	ATA Ile	AGT Ser	AAA Lys 500	ATA Ile	GAA Glu	TTA Leu	CCA Pro	AGT Ser 505	GAA Glu	AAT Asn	ACA Thr	GAA Glu	TCA Ser 510	CTT Leu	ACT Thr	1536 [.]
GAT Asp	TTT Phe	AAT Asn 515	GTA Val	GAT Asp	GTT Val	CCA Pro	GTA Val 520	TAT Tyr	GAA Glu	AAA Lys	CAA Gln	CCC Pro 525	GCT Ala	ATA Ile	AAA Lys	1584
AAA Lys	GTT Val 530	TTT Phe	ACA Thr	GAT Asp	GAA Glu	AAT Asn 535	ACC Thr	ATC Ile	TTT Phe	CAA Gln	TAT Tyr 540	TTA Leu	TAC Tyr	TCT Ser	CAG Gln	1632
ACA Thr 545	Phe	CCT Pro	CTA Leu	AAT Asn	ATA Ile 550	AGA Arg	GAT Asp	ATA Ile	AGT Ser	TTA Leu 555	ACA Thr	TCT Ser	TCA Ser	TTT Phe	GAT Asp 560	1680
GAT Asp	GCA Ala	TTA Leu	TTA Leu	GTT Val 565	TCT Ser	AGC Ser	AAA Lys	GTT Val	TAT Tyr 570	TCA Ser	TTT Phe	TTT Phe	TCT Ser	ATG Met 575	GAT Asp	1728
TAT Tyr	ATT	AAA Lys	ACT Thr 580	GCT Ala	AAT Asn	AAA Lys	GTA Val	GTA Val 585	GAA Glu	GCA Ala	GGA Gly	TTA Leu	TTT Phe 590	GCA Ala	GGT Gly	1776
TGG Trp	GTG Val	AAA Lys 595	CAG Gln	ATA Ile	GTA Val	GAT Asp	GAT Asp 600	TTT Phe	GTA Val	ATC Ile	GAA Glu	GCT Ala 605	AAT Asn	AAA Lys	AGC Ser	1824
AGT Ser	ACT Thr 610	ATG Met	GAT - Asp	AAA Lys	ATT Ile	GCA Ala 615	GAT Asp	ATA Ile	TCT Ser	CTA Leu	ATT Ile 620	GTT Val	CCT Pro	TAT Tyr	ATA Ile	1872
GGA Gly 625	TTA Leu	GCT Ala	TTA Leu	AAT Asn	GTA Val 630	GGA Gly	GAT Asp	GAA Glu	ACA Thr	GCT Ala 635	AAA Lys	GGA Gly	AAT Asn	TTT Phe	GAA Glu 640	1920
AGT Ser			Glu										Phe			1968

GAA Glu	CTI Leu	TTA	ATA Ile 660	Pro	GTA Val	GTI Val	GGA Gly	GTC Val 665	Phe	TTA Leu	Leu	GAA Glu	TCA Ser 670	Tyr	ATT	2016
GAC Asp	AAT Asn	Lys 675	Asn	AAA Lys	ATT	ATT	Lys 680	Thr	ATA Ile	GAT Asp	AAT Asn	GCT Ala 685	TTA Leu	ACT Thr	AAA Lys	2064
AGA Arg	GTG Val 690	Glu	AAA Lys	TGG	ATT Ile	GAT Asp 695	Met	TAC	GGA Gly	TTA Leu	Ile 700	Val	GCG Ala	CAA Gln	TGG Trp	2112
CTC Leu 705	Ser	ACA Thr	GTT Val	AAT Asn	ACT Thr 710	CAA Gln	TTT	TAT Tyr	ACA Thr	ATA Ile 715	AAA Lys	GAG Glu	GGA Gly	ATG Met	TAT Tyr 720	2160
AAG Lys	GCT Ala	TTA Leu	AAT Asn	TAT Tyr 725	CAA Gln	GCA Ala	CAA Gln	Ala	TTG Leu 730	GAA Glu	GAA Glu	ATA Ile	ATA Ile	AAA Lys 735	TAC Tyr	2208
AAA Lys	TAT Tyr	AAT Asn	ATA Ile 740	TAT Tyr	TCT Ser	GAA Glu	GAG Glu	GAA Glu 745	AAG Lys	TCA Ser	AAT Asn	ATT Ile	AAC Asn 750	ATC Ile	AAT Asn	2256
TTT Phe	AAT Asn	GAT Asp 755	ATA Ile	AAT Asn	TCT Ser	AAA Lys	CTT Leu 760	AAT Asn	GAT Asp	GGT Gly	ATT Ile	AAC Asn 765	CAA Gln	GCT Ala	ATG Met	2304
									TGT Cys							2352
AAA Lys 785	AAA Lys	ATG Met	ATT	CCA Pro	TTA Leu 790	GCT Ala	GTA Val	AAA Lys	AAA Lys	TTA Leu 795	CTA Leu	GAC Asp	TTT Phe	GAT Asp	AAT Asn 800	2400
ACT Thr	CTC Leu	AAA Lys	AAA Lys	AAT Asn 805	TTA Leu	TTA Leu	AAT Asn	TAT Tyr	ATA Ile 810	GAT Asp	GAA Glu	AAT Asn	AAA Lys	TTA Leu 815	TAT Tyr	2448
TTA Leu	ATT Ile	GGA Gly	AGT Ser 820	GTA Val	GAA Glu	GAT Asp	GAA Glu	AAA Lys 825	TCA Ser	AAA Lys	GTA Val	GAT Asp	AAA Lys 830	TAC Tyr	TTG Leu	2496
AAA Lys	ACC Thr	ATT Ile 835	ATA Ile	CCA Pro	TTT Phe	GAT Asp	CTT Leu 840	TCA Ser	ACG Thr	TAT Tyr	TCT Ser	AAT Asn 845	ATT Ile	GAA Glu	ATA Ile	2544
									AGC Ser							2592
ATC Ile 865	TTA Leu	AAT Asn	TTA Leu	AGA Arg	TAT Tyr 870	AGA Arg	GAT Asp	AAT Asn	AAT Asn	TTA Leu 875	ATA Ile	GAT Asp	TTA Leu	TCA Ser	GGA Gly 880	2640
									GGG Gly 890				Asn			2688

AAT Asn	CAA Gln	TTT Phe	AAA Lys 900	TTA Leu	ACT Thr	AGT Ser	TCA Ser	GCA Ala 905	Asp	AGT Ser	AAG Lys	ATT Ile	AGA Arg 910	GTC Val	ACT Thr	2736
CAA Gln	AAT Asn	CAG Gln 915	AAT Asn	ATT Ile	ATA Ile	TTT Phe	AAT Asn 920	AGT Ser	ATG Met	TTC Phe	CTT Leu	GAT Asp 925	TTT Phe	AGC Ser	GTT Val	2784
	TTT Phe 930															2832
	ATT Ile															2880
	TGG Trp															2928
	ATA Ile															2976
	GAT Asp							Arg					Thr			3024
	AAT Asn 1010	Leu					Ile					Thr				3072
	ATG Met					Ile					Val					3120
	TTT Phe				Gly					Thr					Met	3168
	TAT Tyr			Ile					Leu					Ile		3216
	ATA Ile		Lys					Ser					Asp			3264
	AAT Asn 1090	Pro					Lys					Phe				3312
	AAA Lys					Lys					Ser					3360
	TTA Leu				Lys					Ser					Tyr	3408

AGA Arg	AAT Asn	TTA Leu	TAT Tyr 114	Ile	GGA Gly	GAA Glu	AAA Lys	TTT Phe 114	Ile	ATA Ile	AGA Arg	AGA Arg	GAG Glu 115	Ser	AAT Asn	3456
TCT Ser	CAA Gln	TCT Ser 115	Ile	AAT Asn	GAT Asp	GAT Asp	ATA Ile 116	Val	AGA Arg	AAA Lys	GAA Glu	GAT Asp 1165	Tyr	ATA Ile	CAT His	3504
CTA Leu	GAT Asp 1170	Leu	GTA Val	CTT Leu	CAC His	CAT His 1175	Glu	GAG Glu	TGG Trp	AGA Arg	GTA Val 1180	Tyr	GCC Ala	TAT Tyr	AAA Lys	3552
TAT Tyr 118	Phe	AAG Lys	GAA Glu	CAG Gln	GAA Glu 1190	Glu	AAA Lys	TTG Leu	TTT Phe	TTA Leu 1195	Ser	ATT Ile	ATA Ile	AGT Ser	GAT Asp 1200	3600
TCT Ser	AAT Asn	GAA Glu	TTT Phe	TAT Tyr 1205	Lys	ACT Thr	ATA Ile	GAA Glu	ATA Ile 1210	AAA Lys)	GAA Glu	TAT Tyr	GAT Asp	GAA Glu 1215	Gln	3648
CCA Pro	TCA Ser	TAT Tyr	AGT Ser 1220	Cys	CAG Gln	TTG Leu	CTT Leu	TTT Phe 1225	Lys	AAA Lys	GAT Asp	Glu	GAA Glu 1230	Ser	ACT Thr	3696
	Asp		Gly					His		TTC Phe	Tyr		Ser			3744
TTA Leu	CGT Arg 1250	Lys	AAG Lys	TAT Tyr	AAA Lys	GAT Asp 1255	Tyr	TTT Phe	TGT Cys	ATA Ile	AGT Ser 1260	Lys	TGG Trp	TAC Tyr	TTA Leu	3792
AAA Lys 1265	Glu	GTA Val	AAA Lys	Arg	AAA Lys 1270	Pro	TAT Tyr	AAG Lys	Ser	AAT Asn 1275	Leu	GGA Gly	TGT Cys	Asn	TGG Trp 1280	3840
			Pro		Asp	GAA Glu		Trp			TAA					3876

- (2) INFORMATION FOR SEQ ID NO:40:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1291 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:40:

Met Pro Val Thr Ile Asn Asn Phe Asn Tyr Asn Asp Pro Ile Asp Asn 1 5 10 15

Asp Asn Ile Ile Met Met Glu Pro Pro Phe Ala Arg Gly Thr Gly Arg

Tyr Tyr Lys Ala Phe Lys Ile Thr Asp Arg Ile Trp Ile Ile Pro Glu 35 40 45

Arg Tyr Thr Phe Gly Tyr Lys Pro Glu Asp Phe Asn Lys Ser Ser Gly 50 60

Ile Phe Asn Arg Asp Val Cys Glu Tyr Tyr Asp Pro Asp Tyr Leu Asn Thr Asn Asp Lys Lys Asn Ile Phe Phe Gln Thr Leu Ile Lys Leu Phe Asn Arg Ile Lys Ser Lys Pro Leu Gly Glu Lys Leu Leu Glu Met Ile 105 Ile Asn Gly Ile Pro Tyr Leu Gly Asp Arg Arg Val Pro Leu Glu Glu Phe Asn Thr Asn Ile Ala Ser Val Thr Val Asn Lys Leu Ile Ser Asn Pro Gly Glu Val Glu Arg Lys Lys Gly Ile Phe Ala Asn Leu Ile Ile Phe Gly Pro Gly Pro Val Leu Asn Glu Asn Glu Thr Ile Asp Ile Gly Ile Gln Asn His Phe Ala Ser Arg Glu Gly Phe Gly Gly Ile Met Gln 185 Met Lys Phe Cys Pro Glu Tyr Val Ser Val Phe Asn Asn Val Gln Glu 200 Asn Lys Gly Ala Ser Ile Phe Asn Arg Arg Gly Tyr Phe Ser Asp Pro 215 Ala Leu Ile Leu Met His Glu Leu Ile His Val Leu His Gly Leu Tyr 230 235 Gly Ile Lys Val Asp Asp Leu Pro Ile Val Pro Asn Glu Lys Lys Phe Phe Met Gln Ser Thr Asp Thr Ile Gln Ala Glu Glu Leu Tyr Thr Phe Gly Gly Gln Asp Pro Ser Ile Ile Ser Pro Ser Thr Asp Lys Ser Ile 280 Tyr Asp Lys Val Leu Gln Asn Phe Arg Gly Ile Val Asp Arg Leu Asn Lys Val Leu Val Cys Ile Ser Asp Pro Asn Ile Asn Ile Asn Ile Tyr 315 Lys Asn Lys Phe Lys Asp Lys Tyr Lys Phe Val Glu Asp Ser Glu Gly Lys Tyr Ser Ile Asp Val Glu Ser Phe Asn Lys Leu Tyr Lys Ser Leu 345 Met Leu Gly Phe Thr Glu Ile Asn Ile Ala Glu Asn Tyr Lys Ile Lys Thr Arg Ala Ser Tyr Phe Ser Asp Ser Leu Pro Pro Val Lys Ile Lys 375 Asn Leu Leu Asp Asn Glu Ile Tyr Thr Ile Glu Glu Gly Phe Asn Ile 395

Ser Asp Lys Asn Met Gly Lys Glu Tyr Arg Gly Gln Asn Lys Ala Ile Asn Lys Gln Ala Tyr Glu Glu Ile Ser Lys Glu His Leu Ala Val Tyr Lys Ile Gln Met Cys Lys Ser Val Lys Val Pro Gly Ile Cys Ile Asp Val Asp Asn Glu Asn Leu Phe Phe Ile Ala Asp Lys Asn Ser Phe Ser 455 Asp Asp Leu Ser Lys Asn Glu Arg Val Glu Tyr Asn Thr Gln Asn Asn Tyr Ile Gly Asn Asp Phe Pro Ile Asn Glu Leu Ile Leu Asp Thr Asp Leu Ile Ser Lys Ile Glu Leu Pro Ser Glu Asn Thr Glu Ser Leu Thr 500 Asp Phe Asn Val Asp Val Pro Val Tyr Glu Lys Gln Pro Ala Ile Lys Lys Val Phe Thr Asp Glu Asn Thr Ile Phe Gln Tyr Leu Tyr Ser Gln Thr Phe Pro Leu Asn Ile Arg Asp Ile Ser Leu Thr Ser Ser Phe Asp 550 Asp Ala Leu Leu Val Ser Ser Lys Val Tyr Ser Phe Phe Ser Met Asp Tyr Ile Lys Thr Ala Asn Lys Val Val Glu Ala Gly Leu Phe Ala Gly - 585 Trp Val Lys Gln Ile Val Asp Asp Phe Val Ile Glu Ala Asn Lys Ser 600 Ser Thr Met Asp Lys Ile Ala Asp Ile Ser Leu Ile Val Pro Tyr Ile Gly Leu Ala Leu Asn Val Gly Asp Glu Thr Ala Lys Gly Asn Phe Glu Ser Ala Phe Glu Ile Ala Gly Ser Ser Ile Leu Leu Glu Phe Ile Pro Glu Leu Leu Ile Pro Val Val Gly Val Phe Leu Leu Glu Ser Tyr Ile 665 Asp Asn Lys Asn Lys Ile Ile Lys Thr Ile Asp Asn Ala Leu Thr Lys Arg Val Glu Lys Trp Ile Asp Met Tyr Gly Leu Ile Val Ala Gln Trp

Leu Ser Thr Val Asn Thr Gln Phe Tyr Thr Ile Lys Glu Gly Met Tyr

Lys Ala Leu Asn Tyr Gln Ala Gln Ala Leu Glu Glu Ile Ile Lys Tyr

725

730

Lys Tyr Asn Ile Tyr Ser Glu Glu Glu Lys Ser Asn Ile Asn Ile Asn 740 745 750

Phe Asn Asp Ile Asn Ser Lys Leu Asn Asp Gly Ile Asn Gln Ala Met 755 760 765

Asp Asn Ile Asn Asp Phe Ile Asn Glu Cys Ser Val Ser Tyr Leu Met 770 780

Lys Lys Met Ile Pro Leu Ala Val Lys Lys Leu Leu Asp Phe Asp Asn 785 790 795 800

Thr Leu Lys Lys Asn Leu Leu Asn Tyr Ile Asp Glu Asn Lys Leu Tyr 805 810 815

Leu Ile Gly Ser Val Glu Asp Glu Lys Ser Lys Val Asp Lys Tyr Leu 820 825 830

Lys Thr Ile Ile Pro Phe Asp Leu Ser Thr Tyr Ser Asn Ile Glu Ile 835 840 845

Leu Ile Lys Ile Phe Asn Lys Tyr Asn Ser Glu Ile Leu Asn Asn Ile 850 855 860

Ile Leu Asn Leu Arg Tyr Arg Asp Asn Asn Leu Ile Asp Leu Ser Gly 865 870 875 880

Tyr Gly Ala Lys Val Glu Val Tyr Asp Gly Val Lys Leu Asn Asp Lys 885 890 895

Asn Gln Phe Lys Leu Thr Ser Ser Ala Asp Ser Lys Ile Arg Val Thr 900 905 910

Gln Asn Gln Asn Ile Ile Phe Asn Ser Met Phe Leu Asp Phe Ser Val 915 920 925

Ser Phe Trp Ile Arg Ile Pro Lys Tyr Arg Asn Asp Asp Ile Gln Asn 930 935 940

Tyr Ile His Asn Glu Tyr Thr Ile Ile Asn Cys Met Lys Asn Asn Ser 945 950 955 960

Gly Trp Lys Ile Ser Ile Arg Gly Asn Arg Ile Ile Trp Thr Leu Ile 965 970 975

Asp Ile Asn Gly Lys Thr Lys Ser Val Phe Phe Glu Tyr Asn Ile Arg 980 985 990

Glu Asp Ile Ser Glu Tyr Ile Asn Arg Trp Phe Phe Val Thr Ile Thr 995 1000 1005

Asn Asn Leu Asp Asn Ala Lys Ile Tyr Ile Asn Gly Thr Leu Glu Ser 1010 1015 1020

Asn Met Asp Ile Lys Asp Ile Gly Glu Val Ile Val Asn Gly Glu Ile 1025 1030 1035 1040

Thr Phe Lys Leu Asp Gly Asp Val Asp Arg Thr Gln Phe Ile Trp Met 1045 1050 1055

Lys Tyr Phe Ser Ile Phe Asn Thr Gln Leu Asn Gln Ser Asn Ile Lys 1060 1065 1070 Glu Ile Tyr Lys Ile Gln Ser Tyr Ser Glu Tyr Leu Lys Asp Phe Trp 1075 1080 1085

Gly Asn Pro Leu Met Tyr Asn Lys Glu Tyr Tyr Met Phe Asn Ala Gly 1090 1095 1100

Asn Lys Asn Ser Tyr Ile Lys Leu Val Lys Asp Ser Ser Val Gly Glu 1105 1110 1115 1120

Ile Leu Ile Arg Ser Lys Tyr Asn Gln Asn Ser Asn Tyr Ile Asn Tyr 1125 1130 1135

Arg Asn Leu Tyr Ile Gly Glu Lys Phe Ile Ile Arg Arg Glu Ser Asn 1140 1145 1150

Ser Gln Ser Ile Asn Asp Asp Ile Val Arg Lys Glu Asp Tyr Ile His 1155 1160 1165

Leu Asp Leu Val Leu His His Glu Glu Trp Arg Val Tyr Ala Tyr Lys 1170 1175 1180

Tyr Phe Lys Glu Glu Glu Glu Lys Leu Phe Leu Ser Ile Ile Ser Asp 1185 1190 1195 1200

Ser Asn Glu Phe Tyr Lys Thr Ile Glu Ile Lys Glu Tyr Asp Glu Gln 1205 1210 1215

Pro Ser Tyr Ser Cys Gln Leu Leu Phe Lys Lys Asp Glu Glu Ser Thr 1220 1225 1230

Asp Asp Ile Gly Leu Ile Gly Ile His Arg Phe Tyr Glu Ser Gly Val 1235 1240 1245

Leu Arg Lys Lys Tyr Lys Asp Tyr Phe Cys Ile Ser Lys Trp Tyr Leu 1250 1255 1260

Lys Glu Val Lys Arg Lys Pro Tyr Lys Ser Asn Leu Gly Cys Asn Trp 1265 1270 1275 1280

Gln Phe Ile Pro Lys Asp Glu Gly Trp Thr Glu 1285 1290

- (2) INFORMATION FOR SEQ ID NO:41:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 3876 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: DNA (genomic)
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 1..3873
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:41:

ATG CCA GTT ACA ATA AAT AAT TTT AAT TAT AAT GAT CCT ATT GAT AAT Met Pro Val Thr Ile Asn Asn Phe Asn Tyr Asn Asp Pro Ile Asp Asn 1 5 10 15

48

AAT Asn	AAT Asn	ATT Ile	ATT Ile 20	ATG Met	ATG Met	GAG Glu	CCT Pro	CCA Pro 25	TTT Phe	GCG Ala	AGA Arg	GGT Gly	ACG Thr 30	GGG Gly	AGA Arg	96
TAT Tyr	TAT Tyr	AAA Lys 35	GCT Ala	TTT Phe	AAA Lys	ATC Ile	ACA Thr 40	GAT Asp	CGT Arg	ATT Ile	TGG Trp	ATA Ile 45	ATA Ile	CCG Pro	GAA Glu	144
AGA Arg	TAT Tyr 50	ACT Thr	TTT Phe	GGA Gly	TAT Tyr	AAA Lys 55	CCT Pro	GAG Glu	GAT Asp	TTT Phe	AAT Asn 60	AAA Lys	AGT Ser	TCC Ser	GGT Gly	192
						TGT Cys										240
ACT Thr	AAT Asn	GAT Asp	AAA Lys	AAG Lys 85	AAT Asn	ATA Ile	TTT Phe	TTA Leu	CAA Gln 90	ACA Thr	ATG Met	ATC Ile	AAG Lys	TTA Leu 95	TTT Phe	288
AAT Asn	AGA Arg	ATC Ile	AAA Lys 100	TCA Ser	AAA Lys	CCA Pro	TTG Leu	GGT Gly 105	GAA Glu	AAG Lys	TTA Leu	TTA Leu	GAG Glu 110	ATG Met	ATT Ile	336
ATA Ile	AAT Asn	GGT Gly 115	ATA Ile	CCT Pro	TAT Tyr	CTT Leu	GGA Gly 120	GAT Asp	AGA Arg	CGT Arg	GTT Val	CCA Pro 125	CTC Leu	GAA Glu	GAG Glu	384
TTT Phe	AAC Asn 130	ACA Thr	AAC Asn	ATT Ile	GCT Ala	AGT Ser 135	GTA Val	ACT Thr	GTT Val	AAT Asn	AAA Lys 140	TTA Leu	ATC Ile	AGT Ser	AAT Asn	432
CCA Pro 145	GGA Gly	GAA Glu	GTG Val	Glu	CGA Arg 150	AAA Lys	AAA Lys	GGT Gly	ATT Ile	TTC Phe 155	GCA Ala	AAT Asn	TTA Leu	ATA Ile	ATA Ile 160	480
TTT Phe	GGA Gly	CCT Pro	GGG Gly	CCA Pro 165	GTT Val	TTA Leu	AAT Asn	GAA Glu	AAT Asn 170	GAG Glu	ACT Thr	ATA Ile	GAT Asp	ATA Ile 175	GGT Gly	528
ATA Ile	CAA Gln	AAT Asn	CAT His 180	TTT Phe	GCA Ala	TCA Ser	AGG Arg	GAA Glu 185	GGC Gly	TTC Phe	GGG Gly	GGT Gly	ATA Ile 190	ATG Met	CAA Gln	576
ATG Met	AAG Lys	TTT Phe 195	TGC Cys	CCA Pro	GAA Glu	TAT Tyr	GTA Val 200	AGC Ser	GTA Val	TTT Phe	AAT Asn	AAT Asn 205	GTT Val	CAA Gln	GAA Glu	624
AAC Asn	AAA Lys 210	GGC Gly	GCA Ala	AGT Ser	ATA Ile	TTT Phe 215	AAT Asn	AGA Arg	CGT Arg	GGA Gly	TAT Tyr 220	TTT Phe	TCA Ser	GAT Asp	CCA Pro	672
GCC Ala 225	TTG Leu	ATA Ile	TTA Leu	ATG Met	CAT His 230	GAA Glu	CTT Leu	ATA Ile	CAT His	GTT Val 235	TTA Leu	CAT His	GGA Gly	TTA Leu	TAT Tyr 240	720
GGC Gly	ATT Ile	AAA Lys	GTA Val	GAT Asp 245	GAT Asp	TTA Leu	CCA Pro	ATT Ile	GTA Val 250	CCA Pro	AAT Asn	GAA Glu	AAA Lys	AAA Lys 255	TTT Phe	768

						GCT Ala									TTT Phe		816
						ATC Ile											864
						AAT Asn 295											912
						TCA Ser											960
						AAA Lys										1	8000
						GAA Glu										1	1056
						ACT Thr										1	104
						AGT Ser 375										1	.152
						ATC Ile										1	.200
						AAA Lys										1	.248
AAT Asn	AAA Lys	CAA Gln	GCT Ala 420	TAT Tyr	GAA Glu	GAA Glu	ATT Ile	AGC Ser 425	AAG Lys	GAG Glu	CAT His	TTG Leu	GCT Ala 430	GTA Val	TAT Tyr	. 1	.296
						AGT Ser										1	.344
						TTC Phe 455										1	.392
						GAA Glu										1	.440
						CCT Pro										1	.488

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TTA Leu	ATA Ile	AGT Ser	AAA Lys 500	ATA Ile	GAA Glu	TTA Leu	CCA Pro	AGT Ser 505	GAA Glu	AAT Asn	ACA Thr	GAA Glu	TCA Ser 510	CTT Leu	ACT Thr	1536
GAT Asp	TTT Phe	AAT Asn 515	GTA Val	GAT Asp	GTT Val	CCA Pro	GTA Val 520	TAT Tyr	GAA Glu	AAA Lys	CAA Gln	CCC Pro 525	GCT Ala	ATA Ile	AAA Lys	1584
AAA Lys	ATT Ile 530	TTT Phe	ACA Thr	GAT Asp	GAA Glu	AAT Asn 535	ACC Thr	ATC Ile	TTT Phe	CAA Gln	TAT Tyr 540	TTA Leu	TAC Tyr	TCT Ser	CAG Gln	1632
						AGA Arg										1680
						AAC Asn										1728
TAT Tyr	ATT Ile	AAA Lys	ACT Thr 580	GCT Ala	AAT Asn	AAA Lys	GTG Val	GTA Val 585	GAA Glu	GCA Ala	GGA Gly	TTA Leu	TTT Phe 590	GCA Ala	GGT Gly	1776
TGG Trp	GTG Val	AAA Lys 595	CAG Gln	ATA Ile	GTA Val	AAT Asn	GAT Asp 600	TTT Phe	GTA Val	ATC Ile	GAA Glu	GCT Ala 605	AAT Asn	AAA Lys	AGC Ser	1824
AAT Asn	ACT Thr 610	ATG Met	GAT Asp	AAA Lys	ATT Ile	GCA Ala 615	GAT Asp	ATA Ile	TCT Ser	CTA Leu	ATT Ile 620	GTT Val	CCT Pro	TAT Tyr	ATA Ile	1872
Gly 625	Leu	Ala	Leu	Asn	Val 630	GGA Gly	Asn	Glu	Thr	Ala 635	Lys	Gly	Asn	Phe	Glu 640	1920
Asn	Ala	Phe	Glu	Ile 645	Ala	GGA Gly	Ala	Ser	Ile 650	Leu	Leu	Glu	Phe	1le 655	Pro	1968
Glu	Leu	Leu	Ile 660	Pro	Val	GTT Val	Gly	Ala 665	Phe	Leu	Leu	Glu	Ser 670	Tyr	Ile	2016
Asp	Asn	Lys 675	Asn	Lys	Ile	ATT	Lys 680	Thr	Ile	Asp	Asn	Ala 685	Leu	Thr	Lys	2064
AGA Arg	AAT Asn 690	GAA Glu	AAA Lys	TGG Trp	AGT Ser	GAT Asp 695	ATG Met	TAC Tyr	GGA Gly	TTA Leu	ATA Ile 700	GTA Val	GCG Ala	CAA Gln	TGG Trp	2112
Leu 705	Ser	Thr	Val	Asn	Thr 710	CAA Gln	Phe	Tyr	Thr	Ile 715	Lys	Glu	Gly	Met	Tyr 720	2160
AAG Lys	GCT Ala	TTA Leu	AAT Asn	TAT Tyr 725	CAA Gln	GCA Ala	CAA Gln	GCA Ala	TTG Leu 730	GAA Glu	GAA Glu	ATA Ile	ATA Ile	AAA Lys 735	TAC Tyr	2208

AGA Arg	TAT Tyr	AAT Asn	ATA Ile 740	TAT Tyr	TCT Ser	GAA Glu	AAA Lys	GAA Glu 745	AAG Lys	TCA Ser	AAT Asn	ATT Ile	AAC Asn 750	ATC Ile	GAT Asp	2256
TTT Phe	AAT Asn	GAT Asp 755	ATA Ile	AAT Asn	TCT Ser	AAA Lys	CTT Leu 760	AAT Asn	GAG Glu	GGT Gly	ATT Ile	AAC Asn 765	CAA Gln	GCT Ala	ATA Ile	2304
					TTT Phe											2352
					TTA Leu 790											2400
					TTG Leu											2448
					GAA Glu											2496
					TTT Phe											2544
					AAT Asn											2592
ATC Ile 865	TTA Leu	AAT Asn	TTA Leu	AGA Arg	TAT Tyr 870	AAG Lys	GAT Asp	AAT Asn	AAT Asn	TTA Leu 875	ATA Ile	GAT Asp	TTA Leu	TCA Ser	GGA Gly 880	2640
TAT Tyr	GGG Gly	GCA Ala	AAG Lys	GTA Val 885	GAG Glu	GTA Val	TAT Tyr	GAT Asp	GGA Gly 890	GTC Val	GAG Glu	CTT Leu	AAT Asn	GAT Asp 895	AAA Lys	2688
					ACT Thr											2736
CAA Gln	AAT Asn	CAG Gln 915	AAT Asn	ATC Ile	ATA Ile	TTT Phe	AAT Asn 920	AGT Ser	GTG Val	TTC Phe	CTT Leu	GAT Asp 925	TTT Phe	AGC Ser	GTT Val	2784
AGC Ser	TTT Phe 930	TGG Trp	ATA Ile	AGA Arg	ATA Ile	CCT Pro 935	AAA Lys	TAT Tyr	AAG Lys	AAT Asn	GAT Asp 940	GGT Gly	ATA Ile	CAA Gln	AAT Asn	2832
TAT Tyr 945	ATT Ile	CAT His	AAT Asn	GAA Glu	TAT Tyr 950	ACA Thr	ATA Ile	ATT Ile	AAT Asn	TGT Cys 955	ATG Met	AAA Lys	AAT Asn	AAT Asn	TCG Ser 960	2880
GGC Gly	TGG Trp	AAA Lys	ATA Ile	TCT Ser 965	ATT Ile	AGG Arg	GGT Gly	AAT Asn	AGG Arg 970	ATA Ile	ATA Ile	TGG Trp	ACT Thr	TTA Leu 975	ATT Ile	2928

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GAT ATA AAT GGA AAA ACC Asp Ile Asn Gly Lys Thr 980	AAA TCG GTA TTT Lys Ser Val Phe 985	TTT GAA TAT AAC Phe Glu Tyr Asn 990	ATA AGA 2976 Ile Arg
GAA GAT ATA TCA GAG TAT Glu Asp Ile Ser Glu Tyr 995			
AAT AAT TTG AAT AAC GCT Asn Asn Leu Asn Asn Ala 1010			
AAT ACA GAT ATT AAA GAT Asn Thr Asp Ile Lys Asp 1025 103	Ile Arg Glu Val	ATT GCT AAT GGT Ile Ala Asn Gly 1035	GAA ATA 3120 Glu Ile 1040
ATA TTT AAA TTA GAT GGT Ile Phe Lys Leu Asp Gly . 1045		Thr Gln Phe Ile	
AAA TAT TTC AGT ATT TTT Lys Tyr Phe Ser Ile Phe 1060			Ile Glu
GAA AGA TAT AAA ATT CAA Glu Arg Tyr Lys Ile Gln 1075			
GGA AAT CCT TTA ATG TAC Gly Asn Pro Leu Met Tyr 1090			
AAT AAA AAT TCA TAT ATT Asn Lys Asn Ser Tyr Ile 1105	Lys Leu Lys Lys		
ATT TTA ACA CGT AGC AAA Ile Leu Thr Arg Ser Lys 1125		Ser Lys Tyr Ile	
AGA GAT TTA TAT ATT GGA Arg Asp Leu Tyr Ile Gly 1140	GAA AAA TTT ATT Glu Lys Phe Ile 1145	ATA AGA AGA AAG Ile Arg Arg Lys 1150	Ser Asn
TCT CAA TCT ATA AAT GAT Ser Gln Ser Ile Asn Asp 1155	GAT ATA GTT AGA Asp Ile Val Arg 1160	AAA GAA GAT TAT Lys Glu Asp Tyr 1165	ATA TAT 3504 Ile Tyr
CTA GAT TTT TTT AAT TTA Leu Asp Phe Phe Asn Leu 1170			
TAT TTT AAG AAA GAG GAA Tyr Phe Lys Lys Glu Glu 1185	Glu Lys Leu Phe	TTA GCT CCT ATA Leu Ala Pro Ile 1195	AGT GAT 3600 Ser Asp 1200
TCT GAT GAG TTT TAC AAT Ser Asp Glu Phe Tyr Asn 1205	ACT ATA CAA ATA Thr Ile Gln Ile 1210	Lys Glu Tyr Asp	GAA CAG 3648 Glu Gln 1215

Pro	ACA Thr	TAT	AGT Ser 1220	Cys	CAG Gln	TTG Leu	CTT Leu	TTT Phe 1225	Lys	AAA Lys	GAT Asp	GAA Glu	GAA Glu 123	Ser	ACT Thr	3696
GAT Asp	GAG Glu	ATA Ile 1235	Gly	TTG Leu	ATT Ile	GGT Gly	ATT Ile 1240	His	CGT Arg	TTC Phe	TAC Tyr	GAA Glu 1245	Ser	GGA Gly	ATT Ile	3744
GTA Val	TTT Phe 1250	Glu	GAG Glu	TAT Tyr	AAA Lys	GAT Asp 1255	Tyr	TTT Phe	TGT Cys	ATA Ile	AGT Ser 1260	Lys	TGG Trp	TAC Tyr	TTA Leu	3792
AAA Lys 126	GAG Glu 5	GTA Val	AAA Lys	AGG Arg	AAA Lys 1270	Pro	TAT Tyr	AAT Asn	TTA Leu	AAA Lys 1275	Leu	GGA Gly	TGT Cys	AAT Asn	TGG Trp 1280	3840
	TTT Phe				Asp					Glu	TAA					3876
(2)	TNEC	ימשפר	PTON	FOD	SEO.	א חד	10 - 42									

(2) INFORMATION FOR SEQ ID NO:42:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1291 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:42:

Met Pro Val Thr Ile Asn Asn Phe Asn Tyr Asn Asp Pro Ile Asp Asn

Asn Asn Ile Ile Met Met Glu Pro Pro Phe Ala Arg Gly Thr Gly Arg

Tyr Tyr Lys Ala Phe Lys Ile Thr Asp Arg Ile Trp Ile Ile Pro Glu

Arg Tyr Thr Phe Gly Tyr Lys Pro Glu Asp Phe Asn Lys Ser Ser Gly

Ile Phe Asn Arg Asp Val Cys Glu Tyr Tyr Asp Pro Asp Tyr Leu Asn 65 70 75 80

Thr Asn Asp Lys Lys Asn Ile Phe Leu Gln Thr Met Ile Lys Leu Phe 90

Asn Arg Ile Lys Ser Lys Pro Leu Gly Glu Lys Leu Leu Glu Met Ile

Ile Asn Gly Ile Pro Tyr Leu Gly Asp Arg Arg Val Pro Leu Glu Glu

Phe Asn Thr Asn Ile Ala Ser Val Thr Val Asn Lys Leu Ile Ser Asn

Pro Gly Glu Val Glu Arg Lys Lys Gly Ile Phe Ala Asn Leu Ile Ile 150

Phe Gly Pro Gly Pro Val Leu Asn Glu Asn Glu Thr Ile Asp Ile Gly Ile Gln Asn His Phe Ala Ser Arg Glu Gly Phe Gly Gly Ile Met Gln 185 Met Lys Phe Cys Pro Glu Tyr Val Ser Val Phe Asn Asn Val Gln Glu 200 Asn Lys Gly Ala Ser Ile Phe Asn Arg Arg Gly Tyr Phe Ser Asp Pro Ala Leu Ile Leu Met His Glu Leu Ile His Val Leu His Gly Leu Tyr Gly Ile Lys Val Asp Asp Leu Pro Ile Val Pro Asn Glu Lys Lys Phe 250 Phe Met Gln Ser Thr Asp Ala Ile Gln Ala Glu Glu Leu Tyr Thr Phe Gly Gly Gln Asp Pro Ser Ile Ile Thr Pro Ser Thr Asp Lys Ser Ile 280 Tyr Asp Lys Val Leu Gln Asn Phe Arg Gly Ile Val Asp Arg Leu Asn 295 Lys Val Leu Val Cys Ile Ser Asp Pro Asn Ile Asn Ile Asn Ile Tyr Lys Asn Lys Phe Lys Asp Lys Tyr Lys Phe Val Glu Asp Ser Glu Gly 330 Lys Tyr Ser Ile Asp Val Glu Ser Phe Asp Lys Leu Tyr Lys Ser Leu 340 Met Phe Gly Phe Thr Glu Thr Asn Ile Ala Glu Asn Tyr Lys Ile Lys Thr Arg Ala Ser Tyr Phe Ser Asp Ser Leu Pro Pro Val Lys Ile Lys 370 Asn Leu Leu Asp Asn Glu Ile Tyr Thr Ile Glu Glu Gly Phe Asn Ile Ser Asp Lys Asp Met Glu Lys Glu Tyr Arg Gly Gln Asn Lys Ala Ile 405 Asn Lys Gln Ala Tyr Glu Glu Ile Ser Lys Glu His Leu Ala Val Tyr Lys Ile Gln Met Cys Lys Ser Val Lys Ala Pro Gly Ile Cys Ile Asp Val Asp Asn Glu Asp Leu Phe Phe Ile Ala Asp Lys Asn Ser Phe Ser 455. 460 Asp Asp Leu Ser Lys Asn Glu Arg Ile Glu Tyr Asn Thr Gln Ser Asn Tyr Ile Glu Asn Asp Phe Pro Ile Asn Glu Leu Ile Leu Asp Thr Asp 485 490

Leu Ile Ser Lys Ile Glu Leu Pro Ser Glu Asn Thr Glu Ser Leu Thr 505 Asp Phe Asn Val Asp Val Pro Val Tyr Glu Lys Gln Pro Ala Ile Lys 520 Lys Ile Phe Thr Asp Glu Asn Thr Ile Phe Gln Tyr Leu Tyr Ser Gln Thr Phe Leu Leu Asp Ile Arg Asp Ile Ser Leu Thr Ser Ser Phe Asp 550 Asp Ala Leu Leu Phe Ser Asn Lys Val Tyr Ser Phe Phe Ser Met Asp Tyr Ile Lys Thr Ala Asn Lys Val Val Glu Ala Gly Leu Phe Ala Gly Trp Val Lys Gln Ile Val Asn Asp Phe Val Ile Glu Ala Asn Lys Ser Asn Thr Met Asp Lys Ile Ala Asp Ile Ser Leu Ile Val Pro Tyr Ile Gly Leu Ala Leu Asn Val Gly Asn Glu Thr Ala Lys Gly Asn Phe Glu Asn Ala Phe Glu Ile Ala Gly Ala Ser Ile Leu Leu Glu Phe Ile Pro 650 Glu Leu Leu Ile Pro Val Val Gly Ala Phe Leu Leu Glu Ser Tyr Ile Asp Asn Lys Asn Lys Ile Ile Lys Thr Ile Asp Asn Ala Leu Thr Lys 680 Arg Asn Glu Lys Trp Ser Asp Met Tyr Gly Leu Ile Val Ala Gln Trp Leu Ser Thr Val Asn Thr Gln Phe Tyr Thr Ile Lys Glu Gly Met Tyr Lys Ala Leu Asn Tyr Gln Ala Gln Ala Leu Glu Glu Ile Ile Lys Tyr 730 Arg Tyr Asn Ile Tyr Ser Glu Lys Glu Lys Ser Asn Ile Asn Ile Asp Phe Asn Asp Ile Asn Ser Lys Leu Asn Glu Gly Ile Asn Gln Ala Ile Asp Asn Ile Asn Asn Phe Ile Asn Gly Cys Ser Val Ser Tyr Leu Met Lys Lys Met Ile Pro Leu Ala Val Glu Lys Leu Leu Asp Phe Asp Asn 790 795 Thr Leu Lys Lys Asn Leu Leu Asn Tyr Ile Asp Glu Asn Lys Leu Tyr

Leu Ile Gly Ser Ala Glu Tyr Glu Lys Ser Lys Val Asn Lys Tyr Leu

Lys Thr Ile Met Pro Phe Asp Leu Ser Ile Tyr Thr Asn Asp Thr Ile 835 840 845

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- Leu Ile Glu Met Phe Asn Lys Tyr Asn Ser Glu Ile Leu Asn Asn Ile 850 855 860
- Ile Leu Asn Leu Arg Tyr Lys Asp Asn Asn Leu Ile Asp Leu Ser Gly 865 870 875 880
- Tyr Gly Ala Lys Val Glu Val Tyr Asp Gly Val Glu Leu Asn Asp Lys 885 890 895
- Asn Gln Phe Lys Leu Thr Ser Ser Ala Asn Ser Lys Ile Arg Val Thr 900 905 910
- Gln Asn Gln Asn Ile Ile Phe Asn Ser Val Phe Leu Asp Phe Ser Val 915 920 925
- Ser Phe Trp Ile Arg Ile Pro Lys Tyr Lys Asn Asp Gly Ile Gln Asn 930 940
- Tyr Ile His Asn Glu Tyr Thr Ile Ile Asn Cys Met Lys Asn Asn Ser 945 950 955 960
- Gly Trp Lys Ile Ser Ile Arg Gly Asn Arg Ile Ile Trp Thr Leu Ile 965 970 975
- Asp Ile Asn Gly Lys Thr Lys Ser Val Phe Phe Glu Tyr Asn Ile Arg 980 985 990
- Glu Asp Ile Ser Glu Tyr Ile Asn Arg Trp Phe Phe Val Thr Ile Thr 995 1000 1005
- Asn Asn Leu Asn Asn Ala Lys Ile Tyr Ile Asn Gly Lys Leu Glu Ser 1010 1015 1020
- Asn Thr Asp Ile Lys Asp Ile Arg Glu Val Ile Ala Asn Gly Glu Ile 1025 1030 1035 1040
- Ile Phe Lys Leu Asp Gly Asp Ile Asp Arg Thr Gln Phe Ile Trp Met 1045 1050 1055
- Lys Tyr Phe Ser Ile Phe Asn Thr Glu Leu Ser Gln Ser Asn Ile Glu 1060 1065 1070
- Glu Arg Tyr Lys Ile Gln Ser Tyr Ser Glu Tyr Leu Lys Asp Phe Trp 1075 1080 1085
- Gly Asn Pro Leu Met Tyr Asn Lys Glu Tyr Tyr Met Phe Asn Ala Gly 1090 1095 1100
- Asn Lys Asn Ser Tyr Ile Lys Leu Lys Lys Asp Ser Pro Val Gly Glu 1105 1115 1120
- Ile Leu Thr Arg Ser Lys Tyr Asn Gln Asn Ser Lys Tyr Ile Asn Tyr 1125 1130 1135
- Arg Asp Leu Tyr Ile Gly Glu Lys Phe Ile Ile Arg Arg Lys Ser Asn 1140 1145 1150
- Ser Gln Ser Ile Asn Asp Asp Ile Val Arg Lys Glu Asp Tyr Ile Tyr 1155 1160 1165

Leu	Asp 117	Phe 0	Phe	Asn	Leu	Asn 117		Glu	Trp	Arg	Val 118		Thr	Tyr	Lys	
Tyr 118		Lys	Lys	Glu	Glu 1190		Lys	Leu	Phe			Pro	Ile	Ser	Asp 1200	
Ser	Asp	Glu	Phe	Tyr 1205		Thr	Ile	Gln	Ile 1210		Glu	Tyr	Asp ·	Glu 121		
Pro	Thr	Tyr	Ser 1220	_	Gln	Leu	Leu	Phe 122	_	Lys	Asp	Glu	Glu 123		Thr	
Asp	Glu	Ile 123		Leu	Ile	Gly	Ile 1240		Arg	Phe	Tyr	Glu 124		Gly	Ile	
Val	Phe 125	Glu 0	Glu	Tyr	Lys	Asp 1255		Phe	Cys	Ile	Ser 1260	_	Trp	Tyr	Leu	
Lys 126		Val	Lys	Arg	Lys 1270		Tyr	Asn	Leu	Lys 1275		Gly	Cys	Asn	Trp 1280	
Gln	Phe	Ile	Pro	Lys 1285		Glu	Gly	Trp	Thr 1290							
(2)	INF	ORMAT	CION	FOR	SEQ	ID N	10:43	3:								
	(i)	(E	A) LE 3) TY C) SI	NGTH PE: RAND	: 15 nucl		ase ació doub	pair I	:s							
	(ii)	MOI (A						clei = "								
	(ix)	-	A) NA	ME/K		CDS 108.	.152	:3								
	(xi)	SEC	UENC	E DE	SCRI	PTIC	N: S	EQ I	D NC	:43:						
AGA:	rctco	GAT C	CCGC	GAAA	T TA	ATAC	GACT	CAC	TATA	GGG	GAAT	TGTO	GAG (GGA1	CAACAA	60
TTC	CCT	CTA G	AAAT	'AATT	T TG	TTTA	ACTI	TAA	GAAG	GAG	ATAT			GGC Gly		116
		CAT His														164
		ATG Met														212
		AAT Asn														260
		AAT Asn														308

GTA Val	TAT	GAT Asp 70	Gly	GTC Val	AAG Lys	CTT Leu	AAT Asn 75	Asp	AAA Lys	AAT Asn	CAA Gln	TTT Phe 80	AAA Lys	TTA Leu	ACT Thr	3	56
AGT Ser	TCA Ser 85	GCA Ala	GAT Asp	AGT Ser	AAG Lys	ATT Ile 90	AGA Arg	GTC Val	ACT Thr	CAA Gln	AAT Asn 95	Gln	AAT Asn	ATT Ile	ATA Ile	4	04
TTT Phe 100	Asn	AGT Ser	ATG Met	TTC Phe	CTT Leu 105	GAT Asp	TTT	AGC Ser	GTT Val	AGC Ser 110	TTT Phe	TGG Trp	ATA Ile	AGG Arg	ATA Ile 115	4	52
CCT Pro	AAA Lys	TAT Tyr	AGG Arg	AAT Asn 120	GAT Asp	GAT Asp	ATA Ile	CAA Gln	AAT Asn 125	TAT Tyr	ATT Ile	CAT His	AAT Asn	GAA Glu 130	TAT Tyr	5	00
ACG Thr	ATA Ile	ATT Ile	AAT Asn 135	Cys	ATG Met	AAA Lys	AAT Asn	AAT Asn 140	TCA Ser	GGC Gly	TGG Trp	AAA Lys	ATA Ile 145	TCT Ser	ATT	5	48
AGG Arg	GGT Gly	AAT Asn 150	AGG Arg	ATA Ile	ATA Ile	TGG Trp	ACC Thr 155	TTA Leu	ATT Ile	GAT Asp	ATA Ile	AAT Asn 160	GGA Gly	AAA Lys	ACC Thr	5:	96
AAA Lys	TCA Ser 165	GTA Val	TTT Phe	TTT Phe	GAA Glu	TAT Tyr 170	AAC Asn	ATA Ile	AGA Arg	GAA Glu	GAT Asp 175	ATA Ile	TCA Ser	GAG Glu	TAT Tyr	64	44
									ACT Thr							65	92
									TCA Ser 205							. 74	40
									ATA Ile							78	88
									ATG Met							83	36
AAT Asn	ACG Thr 245	CAA Gln	TTA Leu	AAT Asn	CAA Gln	TCA Ser 250	AAT Asn	ATT Ile	AAA Lys	GAG Glu	ATA Ile 255	TAT Tyr	AAA Lys	ATT Ile	CAA Gln	88	84
									TGG Trp							93	32
									GGG Gly 285							98	80
									GAA Glu							102	28

			TAT Tyr						1076
			AGA Arg 330						1124
			GAT Asp				 	 	1172
			TAT Tyr						1220
			ATT Ile						1268
			TAT Tyr						1316
			GAA Glu 410						1364
			GAA Glu						1412
			AAA Lys						1460
			GGA Gly	Cys					1508
GGG Gly		TAA							1526

(2) INFORMATION FOR SEQ ID NO:44:

(i) SEQUENCE CHARACTERISTICS:

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- (A) LENGTH: 472 amino acids
 (B) TYPE: amino acid
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:44:

Met Gly His His His His His His His His Ser Ser Gly His

Ile Glu Gly Arg His Met Ala Ser Met Ala Asp Thr Ile Leu Ile Glu

Met Phe Asn Lys Tyr Asn Ser Glu Ile Leu Asn Asn Ile Ile Leu Asn 35 40 45

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Leu Arg Tyr Arg Asp Asn Asn Leu Ile Asp Leu Ser Gly Tyr Gly Ala
50 55 60

Lys Val Glu Val Tyr Asp Gly Val Lys Leu Asn Asp Lys Asn Gln Phe 65 70 75 80

Lys Leu Thr Ser Ser Ala Asp Ser Lys Ile Arg Val Thr Gln Asn Gln 85 90 95

Asn Ile Ile Phe Asn Ser Met Phe Leu Asp Phe Ser Val Ser Phe Trp
100 105 110

Ile Arg Ile Pro Lys Tyr Arg Asn Asp Asp Ile Gln Asn Tyr Ile His 115 120 125

Asn Glu Tyr Thr Ile Ile Asn Cys Met Lys Asn Asn Ser Gly Trp Lys 130 135 140

Ile Ser Ile Arg Gly Asn Arg Ile Ile Trp Thr Leu Ile Asp Ile Asn 145 150 155 160

Gly Lys Thr Lys Ser Val Phe Phe Glu Tyr Asn Ile Arg Glu Asp Ile 165 170 175

Ser Glu Tyr Ile Asn Arg Trp Phe Phe Val Thr Ile Thr Asn Asn Leu 180 185 190

Asp Asn Ala Lys Ile Tyr Ile Asn Gly Thr Leu Glu Ser Asn Met Asp 195 200 205

Ile Lys Asp Ile Gly Glu Val Ile Val Asn Gly Glu Ile Thr Phe Lys 210 225 220

Leu Asp Gly Asp Val Asp Arg Thr Gln Phe Ile Trp Met Lys Tyr Phe 225 230 235

Ser Ile Phe Asn Thr Gln Leu Asn Gln Ser Asn Ile Lys Glu Ile Tyr 245 250 255

Lys Ile Gln Ser Tyr Ser Glu Tyr Leu Lys Asp Phe Trp Gly Asn Pro 260 265 270

Leu Met Tyr Asn Lys Glu Tyr Tyr Met Phe Asn Ala Gly Asn Lys Asn 275 280 285

Ser Tyr Ile Lys Leu Val Lys Asp Ser Ser Val Gly Glu Ile Leu Ile 290 295 300

Arg Ser Lys Tyr Asn Gln Asn Ser Asn Tyr Ile Asn Tyr Arg Asn Leu 305 310 315 320

Tyr Ile Gly Glu Lys Phe Ile Ile Arg Arg Glu Ser Asn Ser Gln Ser 325 330 335

Ile Asn Asp Asp Ile Val Arg Lys Glu Asp Tyr Ile His Leu Asp Leu 340 345 350

Val Leu His His Glu Glu Trp Arg Val Tyr Ala Tyr Lys Tyr Phe Lys 355 360 365

Glu	Gln 370	Glu	Glu	Lys	Leu	Phe 375	Leu	Ser	Ile	Ile	Ser 380	Asp	Ser	Asn	Glu	
Phe 385	Tyr	Lys	Thr	Ile	Glu 390	Ile	Lys	Glu	Tyr	Asp 395	Glu	Gln	Pro	Ser	Tyr 400	
Ser	Cys	Gln	Leu	Leu 405	Phe	Lys	Lys	Asp	Glu 410	Glu	Ser	Thr	Asp	Asp 415	Ile	
Gly	Leu	Ile	Gly 420	Ile	His	Arg	Phe	Tyr 425	Glu	Ser	Gly	Val	Leu 430	Arg	Lys	
Lys	Tyr	Lys 435	Asp	Tyr	Phe	Cys	Ile 440	Ser	Lys	Trp	Tyr	Leu 445	Lys	Glu	Val	
Lys	Arg 450	Lys	Pro	Tyr	Lys	Ser 455	Asn	Leu	Gly	Cys	Asn 460	Trp	Gln	Phe	Ile	
Pro 465	Lys	Asp	Glu	Gly	Trp 470	Thr	Glu									
(2)	INFO	ORMAT	rion	FOR	SEQ	ID N	10:45	5 :								
	(i)	(E	QUENCA) LE B) TY C) ST C) TC	ENGTI (PE : [RANI	i: 15 nucl	647 k Leic ESS:	ase acio doul	pai:	cs							
	(ii)) MOI	LECUI	LE TY	PE:	DNA	(ger	nomi	2)							
	(ix)) FEJ (J	ATURI A) NI B) LO	ME/I	KEY:	CDS	152	23								
	(xi)) SE(OUEN	CE DI	ESCRI	[PTIC	ON: S	SEQ :	ID NO	0:45	:					
AGA'												rtgto	GAG (CGGA	TAACAA	60
TTC	CCT	CTA (gaaa'	raat:	rt to	GTTT?	AACT:	TA	AGAA	GGAG	ATA	racc	ATG Met 1	GGC Gly	CAT	116
CAT	CAT His 5	CAT His	CAT His	CAT His	CAT His	CAT His 10	CAT His	CAC His	AGC Ser	AGC Ser	GGC Gly 15	CAT His	ATC Ile	GAA Glu	GGT Gly	164
CGT Arg 20	CAT His	ATG Met	GCT Ala	AGC Ser	ATG Met 25	GCT Ala	GAT Asp	ACA Thr	ATA Ile	CTA Leu 30	ATA Ile	GAA Glu	ATG Met	TTT Phe	AAT Asn 35	212
AAA Lys	TAT Tyr	AAT Asn	AGC Ser	GAA Glu 40	ATT Ile	TTA Leu	AAT Asn	AAT Asn	ATT Ile 45	ATC Ile	TTA Leu	AAT Asn	TTA Leu	AGA Arg 50	TAT Tyr	260
AAG Lys	GAT Asp	AAT Asn	AAT Asn 55	Leu	ATA Ile	GAT Asp	TTA Leu	TCA Ser 60	GGA Gly	TAT	GGG Gly	GCA Ala	AAG Lys 65	GTA Val	GAG Glu	308

						ATT Ile 90										404
						GAT Asp										452
						GGT Gly										500
						AAA Lys										548
						TGG Trp										596
						TAT Tyr 170										644
ATA Ile 180	AAT Asn	AGA Arg	TGG Trp	TTT Phe	TTT Phe 185	GTA Val	ACT Thr	ATT Ile	ACT Thr	AAT Asn 190	AAT Asn	TTG Leu	AAT Asn	AAC Asn	GCT Ala 195	692
AAA Lys	ATT Ile	TAT Tyr	ATT Ile	AAT Asn 200	GGT Gly	AAG Lys	CTA Leu	GAA Glu	TCA Ser 205	AAT Asn	ACA Thr	GAT Asp	ATT Ile	AAA Lys 210	GAT Asp	740
ATA Ile	AGA Arg	GAA Glu	GTT Val 215	ATT Ile	GCT Ala	AAT Asn	GGT Gly	GAA Glu 220	ATA Ile	ATA Ile	TTT Phe	AAA Lys	TTA Leu 225	GAT Asp	GGT Gly	788
GAT Asp	ATA Ile	GAT Asp 230	AGA Arg	ACA Thr	CAA Gln	TTT Phe	ATT Ile 235	TGG Trp	ATG Met	AAA Lys	TAT Tyr	TTC Phe 240	AGT Ser	ATT	TTT Phe	836
						TCA Ser 250										884
TCA Ser 260	TAT Tyr	AGC Ser	GAA Glu	TAT Tyr	TTA Leu 265	AAA Lys	GAT Asp	TTT Phe	TGG Trp	GGA Gly 270	AAT Asn	CCT Pro	TTA Leu	ATG Met	TAC Tyr 275	932
AAT Asn	AAA Lys	GAA Glu	TAT Tyr	TAT Tyr 280	ATG Met	TTT Phe	AAT Asn	GCG Ala	GGG Gly 285	AAT Asn	AAA Lys	AAT Asn	TCA Ser	TAT Tyr 290	ATT Ile	980
AAA Lys	CTA Leu	AAG Lys	AAA Lys 295	GAT Asp	TCA Ser	CCT Pro	GTA Val	GGT Gly 300	GAA Glu	ATT Ile	TTA Leu	ACA Thr	CGT Arg 305	AGC Ser	AAA Lys	1028
TAT Tyr	AAT Asn	CAA Gln 310	AAT Asn	TCT Ser	AAA Lys	TAT Tyr	ATA Ile 315	AAT Asn	TAT Tyr	AGA Arg	GAT Asp	TTA Leu 320	TAT Tyr	ATT Ile	GGA Gly	1076

GAA Glu	AAA Lys 325	TTT	ATT	ATA Ile	AGA Arg	AGA Arg 330	AAG Lys	TCA Ser	AAT Asn	TCT Ser	CAA Gln 335	TCT Ser	ATA Ile	AAT Asn	GAT Asp	1124
						GAT Asp										1172
						TAT Tyr										1220
						CCT Pro										1268
						TAT Tyr										1316
TTG Leu	CTT Leu 405	TTT Phe	AAA Lys	AAA Lys	GAT Asp	GAA Glu 410	GAA Glu	AGT Ser	ACT Thr	GAT Asp	GAG Glu 415	ATA Ile	GGA Gly	TTG Leu	ATT Ile	1364
GGT Gly 420	ATT Ile	CAT His	CGT Arg	TTC Phe	TAC Tyr 425	GAA Glu	TCT Ser	GGA Gly	ATT Ile	GTA Val 430	TTT Phe	GAA Glu	GAG Glu	TAT Tyr	AAA Lys 435	1412
GAT Asp	TAT Tyr	TTT Phe	TGT Cys	ATA Ile 440	AGT Ser	AAA Lys	TGG Trp	TAC Tyr	TTA Leu 445	AAA Lys	GAG Glu	GTA Val	AAA Lys	AGG Arg 450	AAA Lys	1460
CCA Pro	TAT Tyr	AAT Asn	TTA Leu 455	AAA Lys	TTG Leu	GGA Gly	TGT Cys	AAT Asn 460	TGG Trp	CAG Gln	TTT Phe	ATT Ile	CCT Pro 465	AAA Lys	GAT Asp	1508
		TGG Trp 470			TAA	AGCT	TG C	CGGC	GCA	CT CC	GAG					1547

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(2) INFORMATION FOR SEQ ID NO:46:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 472 amino acids
 (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:46:

Met Gly His His His His His His His His His Ser Ser Gly His

Ile Glu Gly Arg His Met Ala Ser Met Ala Asp Thr Ile Leu Ile Glu

Met Phe Asn Lys Tyr Asn Ser Glu Ile Leu Asn Asn Ile Ile Leu Asn

Leu Arg Tyr Lys Asp Asn Asn Leu Ile Asp Leu Ser Gly Tyr Gly Ala

Lys Val Glu Val Tyr Asp Gly Val Glu Leu Asn Asp Lys Asn Gln Phe Lys Leu Thr Ser Ser Ala Asn Ser Lys Ile Arg Val Thr Gln Asn Gln Asn Ile Ile Phe Asn Ser Val Phe Leu Asp Phe Ser Val Ser Phe Trp 105 Ile Arg Ile Pro Lys Tyr Lys Asn Asp Gly Ile Gln Asn Tyr Ile His Asn Glu Tyr Thr Ile Ile Asn Cys Met Lys Asn Asn Ser Gly Trp Lys Ile Ser Ile Arg Gly Asn Arg Ile Ile Trp Thr Leu Ile Asp Ile Asn 150 Gly Lys Thr Lys Ser Val Phe Phe Glu Tyr Asn Ile Arg Glu Asp Ile Ser Glu Tyr Ile Asn Arg Trp Phe Phe Val Thr Ile Thr Asn Asn Leu Asn Asn Ala Lys Ile Tyr Ile Asn Gly Lys Leu Glu Ser Asn Thr Asp 200 Ile Lys Asp Ile Arg Glu Val Ile Ala Asn Gly Glu Ile Ile Phe Lys Leu Asp Gly Asp Ile Asp Arg Thr Gln Phe Ile Trp Met Lys Tyr Phe Ser Ile Phe Asn Thr Glu Leu Ser Gln Ser Asn Ile Glu Glu Arg Tyr 250 Lys Ile Gln Ser Tyr Ser Glu Tyr Leu Lys Asp Phe Trp Gly Asn Pro Leu Met Tyr Asn Lys Glu Tyr Tyr Met Phe Asn Ala Gly Asn Lys Asn 280 Ser Tyr Ile Lys Leu Lys Lys Asp Ser Pro Val Gly Glu Ile Leu Thr Arg Ser Lys Tyr Asn Gln Asn Ser Lys Tyr Ile Asn Tyr Arg Asp Leu 315 310 Tyr Ile Gly Glu Lys Phe Ile Ile Arg Arg Lys Ser Asn Ser Gln Ser Ile Asn Asp Asp Ile Val Arg Lys Glu Asp Tyr Ile Tyr Leu Asp Phe Phe Asn Leu Asn Gln Glu Trp Arg Val Tyr Thr Tyr Lys Tyr Phe Lys Lys Glu Glu Glu Lys Leu Phe Leu Ala Pro Ile Ser Asp Ser Asp Glu Phe Tyr Asn Thr Ile Gln Ile Lys Glu Tyr Asp Glu Gln Pro Thr Tyr 390 395

Ser	Cys	Gln	Leu	Leu 405	Phe	Lys	Lys	Asp	Glu 410	Glu	Ser	Thr	Asp	Glu 415	Ile	
Gly	Leu	Ile	Gly 420	Ile	His	Arg	Phe	Tyr 425	Glu	Ser	Gly	Ile	Val 430	Phe	Glu	
Glu	Tyr	Lys 435	Asp	Tyr	Phe	Cys	Ile 440	Ser	Lys	Trp	Tyr	Leu 445	Lys	Glu	Val	
Lys	Arg 450	Lys	Pro	Tyr	Asn	Leu 455	Lys	Leu	Gly	Суз	Asn 460	Trp	Gln	Phe	Ile	
Pro 465	Lys	Asp	Glu	Gly	Trp 470	Thr	Glu									
(2)	INFO	RMAT	CION	FOR	SEQ	ID N	10:47	7:								
		(A) (C) (D) (D)	A) LE B) TY C) ST D) TO LECUI	ENGTH (PE: TRANI)POLO	H: 31 nucl DEDNE DGY:	TERI bas eic SS: line othe	e pa acid sing ar	irs l gle iclei								
	(xi)	SEC	UENC	E DE	ESCRI	PTIC	พ: ร	EQ I	D NC	:47:						
CGCC	ATGG	CT G	SATAC	CAATA	C TA	ATAG	TAAA	G								31
(2)	INFO	RMAT	CION	FOR	SEQ	ID N	iO:48) :								
	(i)	(E) (C)	L) LE 3) TY C) ST	NGTH PE: RANE	i: 29 nucl EDNE	TERI bas eic SS: line	e pa acid sing	irs l			•					
	(ii)					othe N:/										
	(xi)	SEC	UENC	E DE	SCRI	PTIC	N: S	EQ I	D NC	:48:						
GCAA	GCTT	TT A	TTCA	GTCC	A CC	CTTC	ATC									29
(2)	INFO	RMAT	CION	FOR	SEQ	ID N	0:49):								
	(i)	(A (B (C	L) LE S) TY C) ST	NGTH PE: RANE	: 37 nucl EDNE	TERI 53 b eic SS: line	ase acid doub	pair l	:s							
	(ii)	MOL	ECUL	E TY	PE:	DNA	(gen	omic	:)							
	(ix)		AN (ME/K	EY:	CDS	750									

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:49:

ATG Met 1	CCA Pro	ACA Thr	ATT Ile	AAT Asn 5	AGT Ser	TTT Phe	AAT Asn	TAT Tyr	AAT Asn 10	GAT Asp	CCT Pro	GTT Val	AAT Asn	AAT Asn 15	AGA Arg	48
						CCA Pro										96
TTT Phe	AAT Asn	ATT Ile 35	ATG Met	AAA Lys	AAT Asn	ATT	TGG Trp 40	ATA Ile	ATT Ile	CCA Pro	GAG Glu	AGA Arg 45	AAT Asn	GTA Val	ATT Ile	144
						TTT Phe 55										192
GAT Asp 65	AGT Ser	AGT Ser	TAT Tyr	TAT Tyr	GAC Asp 70	CCT Pro	AAT Asn	TAT Tyr	TTA Leu	CAA Gln 75	AGT Ser	GAT Asp	CAA Gln	GAA Glu	AAG Lys 80	240
GAT Asp	AAA Lys	TTT Phe	TTA Leu	AAA Lys 85	Ile	GTC Val	ACA Thr	AAA Lys	ATA Ile 90	TTT Phe	AAT Asn	AGA Arg	ATA Ile	AAT Asn 95	GAT Asp	288
AAT Asn	CTT Leu	TCA Ser	GGA Gly 100	AGG Arg	ATT Ile	TTA Leu	TTA Leu	GAA Glu 105	GAA Glu	CTG Leu	TCA Ser	AAA Lys	GCT Ala 110	AAT Asn	CCA Pro	336
TAT Tyr	TTA Leu	GGA Gly 115	Asn	GAT Asp	AAT Asn	ACT Thr	CCA Pro 120	GAT Asp	GGT Gly	GAC Asp	TTC Phe	ATT Ile 125	ATT	AAT Asn	GAT Asp	384
GCA Ala	TCA Ser 130	GCA Ala	GTT Val	CCA Pro	ATT Ile	CAA Gln 135	TTC Phe	TCA Ser	AAT Asn	GGT Gly	AGC Ser 140	CAA Gln	AGC Ser	ATA Ile	CTA Leu	432
TTA Leu 145	CCT Pro	AAT Asn	GTT Val	ATT Ile	ATA Ile 150	ATG Met	GGA Gly	GCA Ala	GAG Glu	CCT Pro 155	GAT Asp	TTA Leu	TTT Phe	GAA Glu	ACT Thr 160	480
AAC Asn	AGT Ser	TCC Ser	AAT Asn	ATT Ile 165	TCT Ser	CTA Leu	AGA Arg	AAT Asn	AAT Asn 170	TAT Tyr	ATG Met	CCA Pro	AGC Ser	AAT Asn 175	CAC His	528
GGT Gly	TTT Phe	GGA Gly	TCA Ser 180	ATA Ile	GCT Ala	ATA Ile	GTA Val	ACA Thr 185	TTC Phe	TCA Ser	CCT Pro	GAA Glu	TAT Tyr 190	TCT Ser	TTT Phe	576
AGA Arg	TTT Phe	AAA Lys 195	GAT Asp	AAT Asn	AGT Ser	ATG Met	AAT Asn 200	GAA Glu	TTT Phe	ATT Ile	CAA Gln	GAT Asp 205	CCT Pro	GCT Ala	CTT Leu	624
ACA Thr	TTA Leu 210	ATG Met	CAT His	GAA Glu	TTA Leu	ATA Ile 215	CAT His	TCA Ser	TTA Leu	CAT His	GGA Gly 220	CTA Leu	TAT Tyr	GGG Gly	GCT Ala	672
AAA Lys 225	GGG Gly	ATT Ile	ACT Thr	ACA Thr	AAG Lys 230	TAT Tyr	ACT Thr	ATA Ile	ACA Thr	CAA Gln 235	AAA Lys	CAA Gln	AAT Asn	CCC	CTA Leu 240	720

					GGT Gly											768
					ATT Ile											816
					GAT Asp											864
					CCA Pro											912
					GAT Asp 310											960
ATA Ile	AAC Asn	AAA Lys	TTT Phe	AAT Asn 325	GAT Asp	ATT Ile	TTT Phe	AAA Lys	AAA Lys 330	TTA Leu	TAC Tyr	AGC Ser	TTT Phe	ACG Thr 335	GAA Glu	1008
TTT Phe	GAT Asp	TTA Leu	GCA Ala 340	ACT Thr	AAA Lys	TTT Phe	CAA Gln	GTT Val 345	AAA Lys	TGT Cys	AGG Arg	CAA Gln	ACT Thr 350	TAT Tyr	ATT Ile	1056
GGA Gly	CAG Gln	TAT Tyr 355	AAA Lys	TAC Tyr	TTC Phe	AAA Lys	CTT Leu 360	TCA Ser	AAC Asn	TTG Leu	TTA Leu	AAT Asn 365	GAT Asp	TCT Ser	ATT Ile	1104
TAT Tyr	AAT Asn 370	ATA Ile	TCA Ser	GAA Glu	GGC Gly	TAT Tyr 375	AAT Asn	ATA Ile	AAT Asn	AAT Asn	TTA Leu 380	AAG Lys	GTA Val	AAT Asn	TTT Phe	1152
AGA Arg 385	GGA Gly	CAG Gln	AAT Asn	GCA Ala	AAT Asn 390	TTA Leu	AAT Asn	CCT Pro	AGA Arg	ATT Ile 395	ATT Ile	ACA Thr	CCA Pro	ATT Ile	ACA Thr 400	 1200
GGT Gly	AGA Arg	GGA Gly	CTA Leu	GTA Val 405	AAA Lys	AAA Lys	ATC Ile	ATT Ile	AGA Arg 410	TTT Phe	TGT Cys	AAA Lys	AAT Asn	ATT Ile 415	GTT Val	1248
TCT Ser	GTA Val	AAA Lys	GGC Gly 420	ATA Ile	AGG Arg	AAA Lys	TCA Ser	ATA Ile 425	TGT Cys	ATC Ile	GAA Glu	ATA Ile	AAT Asn 430	AAT Asn	GGT Gly	1296
GAG Glu	TTA Leu	TTT Phe 435	TTT Phe	GTG Val	GCT Ala	TCC Ser	GAG Glu 440	AAT Asn	AGT Ser	TAT Tyr	AAT Asn	GAT Asp 445	GAT Asp	AAT Asn	ATA Ile	1344
AAT Asn	ACT Thr 450	CCT Pro	AAA Lys	GAA Glu	ATT Ile	GAC Asp 455	GAT Asp	ACA Thr	GTA Val	ACT Thr	TCA Ser 460	AAT Asn	AAT Asn	AAT Asn	TAT Tyr	1392
GAA Glu 465	AAT Asn	GAT Asp	TTA Leu	GAT Asp	CAG Gln 470	GTT Val	ATT Ile	TTA Leu	AAT Asn	TTT Phe 475	AAT Asn	AGT Ser	GAA Glu	TCA Ser	GCA Ala 480	1440

CCT Pro	GGA Gly	CTT Leu	TCA Ser	GAT Asp 485	GAA Glu	AAA Lys	TTA Leu	AAT Asn	TTA Leu 490	ACT Thr	ATC Ile	CAA Gln	AAT Asn	GAT Asp 495	GCT Ala	1	1488
TAT Tyr	ATA Ile	CCA Pro	AAA Lys 500	TAT Tyr	GAT Asp	TCT Ser	AAT Asn	GGA Gly 505	ACA Thr	AGT Ser	GAT Asp	ATA Ile	GAA Glu 510	CAA Gln	CAT His	1	1536
GAT Asp	GTT Val	AAT Asn 515	GAA Glu	CTT Leu	AAT Asn	GTA Val	TTT Phe 520	TTC Phe	TAT Tyr	TTA Leu	GAT Asp	GCA Ala 525	CAG Gln	AAA Lys	GTG Val	1	1584
CCC Pro	GAA Glu 530	GGT Gly	GAA Glu	AAT Asn	AAT Asn	GTC Val 535	AAT Asn	CTC Leu	ACC Thr	TCT Ser	TCA Ser 540	ATT Ile	GAT Asp	ACA Thr	GCA Ala	1	1632
TTA Leu 545	TTA Leu	GAA Glu	CAA Gln	CCT Pro	AAA Lys 550	ATA Ile	TAT Tyr	ACA Thr	TTT Phe	TTT Phe 555	TCA Ser	TCA Ser	GAA Glu	TTT Phe	ATT Ile 560		1680
AAT Asn	AAT Asn	GTC Val	AAT Asn	AAA Lys 565	CCT Pro	GTG Val	CAA Gln	GCA Ala	GCA Ala 570	Leu	TTT Phe	GTA Val	AGC Ser	TGG Trp 575	ATA Ile	1	L728
CAA Gln	CAA Gln	GTA Val	TTA Leu 580	GTA Val	GAT Asp	TTT Phe	ACT Thr	ACT Thr 585	GAA Glu	GCT Ala	AAC Asn	CAA Gln	AAA Lys 590	AGT Ser	ACT Thr	•)(•	1776
GTT Val	GAT Asp	AAA Lys 595	ATT Ile	GCA Ala	GAT Asp	ATT Ile	TCT Ser 600	ATA Ile	GTT Val	GTT Val	CCA Pro	TAT Tyr 605	ATA Ile	GGT Gly	CTT Leu	1	1824
GCT Ala	TTA Leu 610	AAT Asn	ATA Ile	GGA Gly	AAT Asn	GAA Glu 615	GCA Ala	CAA Gln	AAA Lys	GGA Gly	AAT Asn 620	TTT Phe	AAA Lys	GAT Asp	GCA Ala	1	1872
CTT Leu 625	GAA Glu	TTA Leu	TTA Leu	GGA Gly	GCA Ala 630	GGT Gly	ATT Ile	TTA Leu	TTA Leu	GAA Glu 635	TTT Phe	GAA Glu	CCC Pro	Glu	CTT Leu 640	1	1920
TTA Leu	ATT Ile	CCT Pro	ACA Thr	ATT Ile 645	TTA Leu	GTA Val	TTC Phe	ACG Thr	ATA Ile 650	AAA Lys	TCT Ser	TTT Phe	TTA Leu	GGT Gly 655	TCA Ser	1	1968
TCT Ser	GAT Asp	AAT Asn	AAA Lys 660	AAT Asn	AAA Lys	GTT Val	ATT Ile	AAA Lys 665	GCA Ala	ATA Ile	AAT Asn	AAT Asn	GCA Ala 670	TTG Leu	AAA Lys	7	2016
GAA Glu	AGA Arg	GAT Asp 675	GAA Glu	AAA Lys	TGG Trp	AAA Lys	GAA Glu 680	GTA Val	TAT Tyr	AGT Ser	TTT Phe	ATA Ile 685	GTA Val	TCG Ser	AAT Asn	2	2064
TGG Trp	ATG Met 690	ACT Thr	AAA Lys	ATT Ile	AAT Asn	ACA Thr 695	CAA Gln	TTT Phe	AAT Asn	AAA Lys	AGA Arg 700	AAA Lys	GAA Glu	CAA Gln	ATG Met	:	2112
TAT Tyr 705	Gln	GCT Ala	TTA Leu	CAA Gln	AAT Asn 710	CAA Gln	GTA Val	AAT Asn	GCA Ala	CTT Leu 715	AAA Lys	GCA Ala	ATA Ile	ATA Ile	GAA Glu 720	:	2160

TCI Ser	' AAG ' Lys	TAT	AAT Asn	AGT Ser 725	TAT	ACT Thr	TTA Leu	GAA Glu	GAA Glu 730	AAA Lys	AAT Asn	GAG Glu	CTT Leu	ACA Thr 735	AAT Asn		2208
AAA Lys	TAT Tyr	GAT Asp	ATT Ile 740	GAG Glu	CAA Gln	ATA Ile	GAA Glu	AAT Asn 745	GAA Glu	CTT Leu	AAT Asn	CAA Gln	AAG Lys 750	GTT Val	TCT Ser		2256
	GCA Ala		Asn														2304
TAT Tyr	TTA Leu 770	ATG Met	AAA Lys	TTA Leu	ATA Ile	AAT Asn 775	GAA Glu	GTA Val	AAA Lys	ATT Ile	AAT Asn 780	AAA Lys	TTA Leu	AGA Arg	GAA Glu		2352
	GAT Asp															•	2400
	TCA Ser																2448
	ACC Thr																2496
	AAA Lys																2544
	AGT Ser 850																2592
	TCA Ser																2640
	CCA Pro																2688
	GTT Val																2736
	AAT Asn																2784
	ATA Ile 930														AGG . Arg		2832
	AAT Asn																2880

	Asn Ser Gly Ile	AAT CAA AAA TTA GCA Asn Gln Lys Leu Ala 970	
TAT GGT AAC GCA AAT Tyr Gly Asn Ala Asn 980	GGT ATT TCT GAT Gly Ile Ser Asp 985	TAT ATA AAT AAG TGG Tyr Ile Asn Lys Trp 990	ATT TTT 2976 Ile Phe
		GAT TCT AAA CTT TAT Asp Ser Lys Leu Tyr 1005	
		TTA AAT TTA GGT AAT Leu Asn Leu Gly Asn 1020	
		GTT AAT TGT AGT TAT Val Asn Cys Ser Tyr 1035	
	Tyr Phe Asn Ile	TTT GAT AAA GAA TTA Phe Asp Lys Glu Leu 1050	
		GAA CCT AAT GCA AAT Glu Pro Asn Ala Asn 1070	Ile Leu
		TAT GAC AAA GAA TAC Tyr Asp Lys Glu Tyr 1085	
		ATT AAT AGG AGA ACA Ile Asn Arg Arg Thr 1100	
		ACT ATT CTT TTA GCT Thr Ile Leu Leu Ala 1115	
	Lys Val Lys Ile	CAA AGA GTT AAT AAT Gln Arg Val Asn Asn 1130	
ACT AAC GAT AAT CTT Thr Asn Asp Asn Leu 1140	GTT AGA AAG AAT Val Arg Lys Asn 1145	GAT CAG GTA TAT ATT Asp Gln Val Tyr Ile 1150	Asn Phe
		TTA TAT GCT GAT ACA Leu Tyr Ala Asp Thr 1165	
		TCA TCA TCT GGC AAT Ser Ser Ser Gly Asn 1180	
		GGA TGT ACA ATG AAT Gly Cys Thr Met Asn 1195	

				Asn			TTG Leu		Gly				Thr	3648	
		 	Thr				ACA Thr 1225	His				Thr		3696	
		 Phe					ATT Ile			 	Gly			3744	
GAA Glu	AAA Lys 1250													3753	
(2)	Time	 TON	BOB	000	TD N	10 . 5 .									

- (2) INFORMATION FOR SEQ ID NO:50:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1250 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:50:

Met Pro Thr Ile Asn Ser Phe Asn Tyr Asn Asp Pro Val Asn Asn Arg

1 10 15

Thr Ile Leu Tyr Ile Lys Pro Gly Gly Cys Gln Gln Phe Tyr Lys Ser 20 25 30

Phe Asn Ile Met Lys Asn Ile Trp Ile Ile Pro Glu Arg Asn Val Ile 35 40 45

Gly Thr Ile Pro Gln Asp Phe Leu Pro Pro Thr Ser Leu Lys Asn Gly 50 55 60

Asp Ser Ser Tyr Tyr Asp Pro Asn Tyr Leu Gln Ser Asp Gln Glu Lys
65 70 75 80

Asp Lys Phe Leu Lys Ile Val Thr Lys Ile Phe Asn Arg Ile Asn Asp 85 90 95

Asn Leu Ser Gly Arg Ile Leu Leu Glu Glu Leu Ser Lys Ala Asn Pro 100 105 110

Tyr Leu Gly Asn Asp Asn Thr Pro Asp Gly Asp Phe Ile Ile Asn Asp 115 120 125

Ala Ser Ala Val Pro Ile Gln Phe Ser Asn Gly Ser Gln Ser Ile Leu 130 135 140

Leu Pro Asn Val Ile Ile Met Gly Ala Glu Pro Asp Leu Phe Glu Thr 145 150 155 160

Asn Ser Ser Asn Ile Ser Leu Arg Asn Asn Tyr Met Pro Ser Asn His 165 170 175

Gly Phe Gly Ser Ile Ala Ile Val Thr Phe Ser Pro Glu Tyr Ser Phe 180 185 190 Arg Phe Lys Asp Asn Ser Met Asn Glu Phe Ile Gln Asp Pro Ala Leu 195 200 205

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- Thr Leu Met His Glu Leu Ile His Ser Leu His Gly Leu Tyr Gly Ala 210 215 220
- Lys Gly Ile Thr Thr Lys Tyr Thr Ile Thr Gln Lys Gln Asn Pro Leu 225 230 235 240
- Ile Thr Asn Ile Arg Gly Thr Asn Ile Glu Glu Phe Leu Thr Phe Gly 245 250 255
- Gly Thr Asp Leu Asn Ile Ile Thr Ser Ala Gln Ser Asn Asp Ile Tyr 260 265 270
- Thr Asn Leu Leu Ala Asp Tyr Lys Lys Ile Ala Ser Lys Leu Ser Lys
 275
 280
 285
- Val Gln Val Ser Asn Pro Leu Leu Asn Pro Tyr Lys Asp Val Phe Glu 290 295 300
- Ala Lys Tyr Gly Leu Asp Lys Asp Ala Ser Gly Ile Tyr Ser Val Asn 305 310 315 320
- Ile Asn Lys Phe Asn Asp Ile Phe Lys Lys Leu Tyr Ser Phe Thr Glu 325 330 335
- Phe Asp Leu Ala Thr Lys Phe Gln Val Lys Cys Arg Gln Thr Tyr Ile 340 345 350
- Gly Gln Tyr Lys Tyr Phe Lys Leu Ser Asn Leu Leu Asn Asp Ser Ile 355 360 365
- Tyr Asn Ile Ser Glu Gly Tyr Asn Ile Asn Asn Leu Lys Val Asn Phe 370 375 380
- Arg Gly Gln Asn Ala Asn Leu Asn Pro Arg Ile Ile Thr Pro Ile Thr 385 390 395 400
- Gly Arg Gly Leu Val Lys Lys Ile Ile Arg Phe Cys Lys Asn Ile Val 405 410 415
- Ser Val Lys Gly Ile Arg Lys Ser Ile Cys Ile Glu Ile Asn Asn Gly 420 425 430
- Glu Leu Phe Phe Val Ala Ser Glu Asn Ser Tyr Asn Asp Asp Asn Ile 435 440 445
- Asn Thr Pro Lys Glu Ile Asp Asp Thr Val Thr Ser Asn Asn Asn Tyr 450 455 460
- Glu Asn Asp Leu Asp Gln Val Ile Leu Asn Phe Asn Ser Glu Ser Ala 465 470 475 480
- Pro Gly Leu Ser Asp Glu Lys Leu Asn Leu Thr Ile Gln Asn Asp Ala 485 490 495
- Tyr Ile Pro Lys Tyr Asp Ser Asn Gly Thr Ser Asp Ile Glu Gln His
 500 505 510
- Asp Val Asn Glu Leu Asn Val Phe Phe Tyr Leu Asp Ala Gln Lys Val 515 520 525

	Pro	530	GIY	Glu	Asn	Asn	Va1 535		Leu	Thr	Ser	540		Asp	Thr	Ala
	Leu 545	·Leu	Glu	Gln	Pro	Lys 550	Ile	Tyr	Thr	Phe	Phe 555	Ser	Ser	Glu	Phe	Ile 560
	Asn	Asn	Val	Asn	Lys 565	Pro	Val	Gln	Ala	Ala 570	Leu	Phe	Val	Ser	Trp 575	Ile
	Gln	Gln	Val	Leu 580	Val	Asp	Phe	Thr	Thr 585	Glu	Ala	Asn	Gln	Lys 590	Ser	Thr
	Val	Asp	Lys 595	Ile	Ala	Asp	Ile	Ser 600	Ile	Val	Val	Pro	Tyr 605	Ile	Gly	Leu
	Ala	Leu 610	Asn	Ile	Gly	Asn	Glu 615	Ala	Gln	Lys	Gly	Asn 620	Phe	Lys	Asp	Ala
	Leu 625	Glu	Leu	Leu	Gly	Ala 630	Gly	Ile	Leu	Leu	Glu 635	Phe	Glu	Pro	Glu	Leu 640
	Leu	Ile	Pro	Thr	Ile 645	Leu	Val	Phe	Thr	Ile 650	Lys	Ser	Phe	Leu	Gly 655	Ser
	Ser	Asp	Asn	Lys 660	Asn	Lys	Val	Ile	Lys 665	Ala	Ile	Asn	Asn	Ala 670	Leu	Lys
	Glu	Arg	Asp 675	Glu	Lys	Trp	Lys	Glu 680	Val	Tyr	Ser	Phe	Ile 685	Val	Ser	Asn
	Trp	Met 690	Thr	Lys	Ile	Asn	Thr 695	Gln	Phe	Asn	Lys:	Arg 700	Lys	Glu	Gln	Met
	Tyr 705	Gln	Ala	Leu	Gln	Asn 710	Gln	Val	Asn	Ala	Leu 715	Lys	Ala	Ile	Ile	Glu 720
	Ser	Lys	Tyr	Asn	Ser 725	Tyr	Thr	Leu	Glu	Glu 730	Lys	Asn	Glu	Leu	Thr 735	Asn
	Lys	Tyr	Asp	Ile 740	Glu	Gln	Ile	Glu	Asn 745	Glu	Leu	Asn	Gln	Lys 750	Val	Ser
	Ile	Ala	Met 755	Asn	Asn	Ile	Asp	Arg 760	Phe	Leu	Thr	Glu	Ser 765	Ser	Ile	Ser
	Tyr	Leu 770	Met	Lys	Leu	Ile	Asn 775	Glu	Val	Lys	Ile	Asn 780	Lys	Leu	Arg	Glu
	Tyr 785	Asp	Glu	Asn	Val	Lys 790	Thr	Tyr	Leu	Leu	Asp 795	Tyr	Ile	Ile	Lys	His 800
•	Gly	Ser	Ile	Leu	Gly 805	Glu	Ser	Gln	Gln	Glu 810	Leu	Asn	Ser	Met	Val 815	Ile
•	Asp	Thr	Leu	Asn 820	Asn	Ser	Ile	Pro	Phe 825	Lys	Leu	Ser	Ser	Tyr 830	Thr	Asp
	Asp	Lys	Ile 835	Leu	Ile	Ser	Tyr	Phe 840	Asn	Lys	Phe	Phe	Lys 845	Arg	Ile	Lys
	Ser	Ser 850	Ser	Val	Leu	Asn	Met 855	Arg	Tyr	Lys	Asn	Asp 860	Lys	Tyr	Val	Asp

Thr Ser Gly Tyr Asp Ser Asn Ile Asn Ile Asn Gly Asp Val Tyr Lys 865 870 875

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- Tyr Pro Thr Asn Lys Asn Gln Phe Gly Ile Tyr Asn Asp Lys Leu Ser 885 890 895
- Glu Val Asn Ile Ser Gln Asn Asp Tyr Ile Ile Tyr Asp Asn Lys Tyr 900 905 910
- Lys Asn Phe Ser Ile Ser Phe Trp Val Arg Ile Pro Asn Tyr Asp Asn 915 920 925
- Lys Ile Val Asn Val Asn Asn Glu Tyr Thr Ile Ile Asn Cys Met Arg 930 935 940
- Asp Asn Asn Ser Gly Trp Lys Val Ser Leu Asn His Asn Glu Ile Ile 945 950 955 960
- Trp Thr Leu Gln Asp Asn Ser Gly Ile Asn Gln Lys Leu Ala Phe Asn 965 970 975
- Tyr Gly Asn Ala Asn Gly Ile Ser Asp Tyr Ile Asn Lys Trp Ile Phe 980 985 990
- Val Thr Ile Thr Asn Asp Arg Leu Gly Asp Ser Lys Leu Tyr Ile Asn 995 1000 1005
- Gly Asn Leu Ile Asp Lys Lys Ser Ile Leu Asn Leu Gly Asn Ile His 1010 1015 1020
- Val Ser Asp Asn Ile Leu Phe Lys Ile Val Asn Cys Ser Tyr Thr Arg 1025 1030 1035 1040
- Tyr Ile Gly Ile Arg Tyr Phe Asn Ile Phe Asp Lys Glu Leu Asp Glu 1045 1050 1055
- Thr Glu Ile Gln Thr Leu Tyr Asn Asn Glu Pro Asn Ala Asn Ile Leu 1060 1065 1070
- Lys Asp Phe Trp Gly Asn Tyr Leu Leu Tyr Asp Lys Glu Tyr Tyr Leu 1075 1080 1085
- Leu Asn Val Leu Lys Pro Asn Asn Phe Ile Asn Arg Arg Thr Asp Ser 1090 1095 1100
- Thr Leu Ser Ile Asn Asn Ile Arg Ser Thr Ile Leu Leu Ala Asn Arg 1105 1110 1115 1120
- Leu Tyr Ser Gly Ile Lys Val Lys Ile Gln Arg Val Asn Asn Ser Ser 1125 1130 1135
- Thr Asn Asp Asn Leu Val Arg Lys Asn Asp Gln Val Tyr Ile Asn Phe 1140 1145 1150
- Val Ala Ser Lys Thr His Leu Leu Pro Leu Tyr Ala Asp Thr Ala Thr 1155 1160 1165
- Thr Asn Lys Glu Lys Thr Ile Lys Ile Ser Ser Ser Gly Asn Arg Phe 1170 1180
- Asn Gln Val Val Val Met Asn Ser Val Gly Cys Thr Met Asn Phe Lys 1185 1190 1195 1200

Asn	Asn	Asn	Gly	Asn 120	Asn 5	Ile	Gly	Leu	Leu 121		Phe	Lys	Ala	Asp 121		
Val	Val	Ala	Ser 122		Trp	Tyr	Tyr	Thr 122		Met	Arg	Asp	Asn 123		Asn	
Ser	Asn	Gly 123		Phe	Trp	Asn	Phe 124		Ser	Glu	Glu	His 124	-	Trp	Gln	
Glu	Lys 1250)								•						
(2)	INF	ORMA'	TION	FOR	SEQ	ID I	NO:5	1:								
	(i)	() ()	A) L: B) T C) S'	engti Ype : Irani	HARAG H: 3' nucl DEDNI OGY:	759 l leic ESS:	oase acio doul	pai: i	rs						,	
	(ii)	MO	LECU	LE T	YPE:	DNA	(ger	nomi	c)							
	(ix)	FEZ			KEY:	CDS										
					ION:		3756									
	(xi)	SE	QUEN	CE DI	ESCRI	PTIC	ON: S	SEQ 1	ID NO	0:51	:					
					AGT Ser											48
					AAA Lys											96
					AAT Asn											144
					GAT Asp											192
					GAC Asp 70											240
					ATA Ile											288
					ATT Ile											336
					AAT Asn											384

	GCA Ala	Ser 130	: Ala	GTT Val	GAG Glu	ATT Ile	Lys 135	Phe	TCA Ser	AAT Asn	GGT Gly	AGC Ser 140	Gln	GAC Asp	ATA Ile	CTA Leu		432
	TTA Leu 145	Pro	AAT Asn	GTT Val	ATT Ile	ATA Ile 150	ATG Met	GGA Gly	GCA Ala	GAG Glu	CCT Pro 155	Asp	TTA Leu	TTT	GAA Glu	ACT Thr 160		480
	AAC Asn	AGT Ser	TCC Ser	AAT Asn	ATT Ile 165	Ser	CTA Leu	AGA Arg	AAT Asn	AAT Asn 170	Tyr	ATG Met	CCA Pro	AGC Ser	AAT Asn 175	CAC His		528
	GGT Gly	TTT Phe	GGA Gly	TCA Ser 180	ATA Ile	GCT Ala	ATA Ile	GTA Val	ACA Thr 185	Phe	TCA Ser	CCT Pro	GAA Glu	TAT Tyr 190	TCT Ser	TTT Phe		576
	AGA Arg	TTT Phe	AAT Asn 195	Asp	AAT Asn	AGT Ser	ATG Met	AAT Asn 200	Glu	TTT Phe	ATT Ile	CAA Gln	GAT Asp 205	CCT Pro	GCT Ala	CTT Leu		624
	ACA Thr	TTA Leu 210	ATG Met	CAT His	GAA Glu	TTA Leu	ATA Ile 215	CAT His	TCA Ser	TTA Leu	CAT His	GGA Gly 220	CTA Leu	TAT Tyr	GGG Gly	GCT Ala	(672
	AAA Lys 225	GGG Gly	ATT Ile	ACT Thr	ACA Thr	AAG Lys 230	TAT Tyr	ACT Thr	ATA Ile	ACA Thr	CAA Gln 235	AAA Lys	CAA Gln	AAT Asn	CCC Pro	CTA Leu 240	•	720
	ATA Ile	ACA Thr	AAT Asn	ATA Ile	AGA Arg 245	GGT Gly	ACA Thr	AAT Asn	ATT Ile	GAA Glu 250	GAA Glu	TTC Phe	TTA Leu	ACT Thr	TTT Phe 255	GGA Gly	5	768
	GGT Gly	ACT Thr	GAT Asp	TTA Leu 260	AAC Asn	ATT Ile	ATT Ile	ACT Thr	AGT Ser 265	GCT Ala	CAG Gln	TCC Ser	AAT Asn	GAT Asp 270	ATC Ile	TAT Tyr	8	316
	ACT Thr	AAT Asn	CTT Leu 275	CTA Leu	GCT Ala	GAT Asp	TAT Tyr	AAA Lys 280	AAA Lys	ATA Ile	GCG Ala	TCT Ser	AAA Lys 285	CTT Leu	AGC Ser	AAA Lys	8	864
	GTA Val	CAA Gln 290	GTA Val	TCT Ser	AAT Asn	CCA Pro	CTA Leu 295	CTT Leu	AAT Asn	CCT Pro	TAT Tyr	AAA Lys 300	GAT Asp	GTT Val	TTT Phe	GAA Glu	9	12
													TAT Tyr				9	60
	ATA Ile	AAC Asn	AAA Lys	TTT Phe	AAT Asn 325	GAT Asp	ATT Ile	TTT Phe	AAA Lys	AAA Lys 330	TTA Leu	TAC Tyr	AGC Ser	TTT Phe	ACG Thr 335	GAA Glu	10	80
;	TTT Phe	GAT Asp	TTA Leu	GCA Ala 340	ACT Thr	AAA Lys	TTT Phe	CAA Gln	GTT Val 345	AAA Lys	TGT Cys	AGG Arg	CAA Gln	ACT Thr 350	TAT Tyr	ATT Ile	10	56
(GGA Gly	CAG Gln	TAT Tyr 355	AAA Lys	TAC Tyr	TTC Phe	Lys	CTT Leu 360	TCA Ser	AAC Asn	TTG Leu	TTA Leu	AAT Asn 365	GAT Asp	TCT Ser	ATT Ile	11	04

TA1 Tyr	AAT Asn 370	lle	TCA Ser	GAA Glu	GGC Gly	TAT Tyr 375	Asn	TATA	AAT Asn	' AAT Asn	TTA Leu 380	Lys	GTA Val	AAT Asn	TTT		1152
AGA Arg 385	Gly	CAG Gln	AAT Asn	GCA Ala	AAT Asn 390	Leu	AAT Asn	CCT Pro	AGA Arg	ATT Ile 395	Ile	ACA Thr	CCA Pro	ATT	ACA Thr 400		1200
GGT Gly	AGA Arg	GGA Gly	CTA Leu	GTA Val 405	Lys	AAA Lys	ATC Ile	ATT	AGA Arg 410	Phe	TGT Cys	AAA Lys	AAT Asn	ATT Ile 415	GTT Val		1248
TCT Ser	GTA Val	AAA Lys	GGC Gly 420	ATA Ile	AGG Arg	AAA Lys	TCA Ser	ATA Ile 425	TGT Cys	ATC Ile	GAA Glu	ATA Ile	AAT Asn 430	AAT Asn	GGT Gly		1296
GAG Glu	TTA Leu	TTT Phe 435	TTT	GTG Val	GCT Ala	TCC Ser	GAG Glu 440	AAT Asn	AGT Ser	TAT Tyr	AAT Asn	GAT Asp 445	GAT Asp	AAT Asn	ATA Ile		1344
AAT Asn	ACT Thr 450	CCT Pro	AAA Lys	GAA Glu	ATT Ile	GAC Asp 455	GAT Asp	ACA Thr	GTA Val	ACT Thr	TCA Ser 460	AAT Asn	AAT Asn	AAT Asn	TAT Tyr		1392
GAA Glu 465	AAT Asn	GAT Asp	TTA Leu	GAT Asp	CAG Gln 470	GTT Val	ATT Ile	TTA Leu	AAT Asn	TTT Phe 475	AAT Asn	AGT Ser	GAA Glu	TCA Ser	GCA Ala 480		1440
CCT Pro	GGA Gly	CTT Leu	Ser	GAT Asp 485	GAA Glu	AAA Lys	TTA Leu	AAT Asn	TTA Leu 490	ACT Thr	ATC Ile	CAA Gln	AAT Asn	GAT Asp 495	GCT Ala		1488
TAT Tyr	ATA Ile	CCA Pro	AAA Lys 500	TAT Tyr	GAT Asp	TCT Ser	AAT Asn	GGA Gly 505	ACA Thr	AGT Ser	GAT Asp	ATA Ile	GAA Glu 510	CAA Gln	CAT His		1536
GAT Asp	GTT Val	AAT Asn 515	GAA Glu	CTT Leu	AAT Asn	GTA Val	TTT Phe 520	TTC Phe	TAT Tyr	TTA Leu	GAT Asp	GCA Ala 525	CAG Gln	AAA Lys	GTG Val		1584
CCC Pro	GAA Glu 530	GGT Gly	GAA Glu	AAT Asn	AAT Asn	GTC Val 535	AAT Asn	CTC Leu	ACC Thr	TCT Ser	TCA Ser 540	ATT Ile	GAT Asp	ACA Thr	GCA Ala		1632
TTA Leu 545	TTA Leu	GAA Glu	CAA Gln	CCT Pro	AAA Lys 550	ATA Ile	TAT Tyr	ACA Thr	TTT Phe	TTT Phe 555	TCA Ser	TCA Ser	GAA Glu	TTT Phe	ATT Ile 560	,	1680
AAT Asn	AAT Asn	GTC Val	AAT Asn	AAA Lys 565	CCT Pro	GTG Val	CAA Gln	GCA Ala	GCA Ala 570	TTA Leu	TTT Phe	GTA Val	AGC Ser	TGG Trp 575	ATA Ile		1728
CAA Gln	CAA . Gln	GTG Val	TTA Leu 580	GTA Val	GAT Asp	TTT Phe	ACT Thr	ACT Thr 585	GAA Glu	GCT Ala	AAC Asn	CAA Gln	AAA Lys 590	AGT Ser	ACT Thr		1776
GTT Val	GAT Asp	AAA Lys 595	ATT Ile	GCA Ala	GAT Asp	Ile	TCT Ser 600	ATA Ile	GTT Val	GTT Val	CCA Pro	TAT Tyr 605	ATA Ile	GGT Gly	CTT Leu		1824

GCT Ala	TTA Leu 610	Asn	ATA Ile	GGA Gly	AAT Asn	GAA Glu 615	Ala	CAA Gln	AAA Lys	GGA Gly	AAT Asn 620	Phe	AAA Lys	GAT Asp	GCA Ala	1872
CTT Leu 625	Glu	TTA Leu	TTA Leu	GGA Gly	GCA Ala 630	Gly	ATT Ile	TTA Leu	TTA Leu	GAA Glu 635	Phe	GAA Glu	CCC Pro	GAG Glu	CTT Leu 640	1920
TTA Leu	ATT Ile	CCT Pro	ACA Thr	ATT Ile 645	TTA Leu	GTA Val	TTC Phe	ACG Thr	ATA Ile 650	AAA Lys	TCT Ser	TTT Phe	TTA Leu	GGT Gly 655	TCA Ser	1968
TCT Ser	GAT Asp	AAT Asn	AAA Lys 660	AAT Asn	AAA Lys	GTT Val	ATT Ile	AAA Lys 665	GCA Ala	ATA Ile	AAT Asn	AAT Asn	GCA Ala 670	TTG Leu	AAA Lys	2016
GAA Glu	AGA Arg	GAT Asp 675	GAA Glu	AAA Lys	TGG Trp	AAA Lys	GAA Glu 680	GTA Val	TAT Tyr	AGT Ser	TTT Phe	ATA Ile 685	GTA Val	TCG Ser	AAT Asn	2064
TGG Trp	ATG Met 690	ACT Thr	AAA Lys	ATT Ile	AAT Asn	ACA Thr 695	CAA Gln	TTT Phe	AAT Asn	AAA Lys	AGA Arg 700	AAA Lys	GAA Glu	CAA Gln	ATG Met	2112
TAT Tyr 705	CAA Gln	GCT Ala	TTA Leu	CAA Gln	AAT Asn 710	CAA Gln	GTA Val	AAT Asn	GCA Ala	ATT Ile 715	AAA Lys	ACA Thr	ATA Ile	ATA Ile	GAA Glu 720	2160
TCT Ser	AAG Lys	TAT Tyr	AAT Asn	AGT Ser 725	TAT Tyr	ACT Thr	TTA Leu	GAG Glu	GAA Glu 730	AAA Lys	AAT Asn	GAG Glu	CTT Leu	ACA Thr 735	AAT Asn	2208
AAA Lys	TAT Tyr	GAT Asp	ATT Ile 740	AAG Lys	CAA Gln	ATA Ile	GAA Glu	AAT Asn 745	GAA Glu	CTT Leu	AAT Asn	CAA Gln	AAG Lys 750	GTT Val	TCT Ser	2256
		ATG Met 755														2304
		ATG Met														2352
TAT Tyr 785	GAT Asp	GAG Glu	AAT Asn	GTC Val	AAA Lys 790	ACG Thr	TAT Tyr	TTA Leu	TTG Leu	AAT Asn 795	TAT Tyr	ATT Ile	ATA Ile	CAA Gln	CAT His 800	2400
		ATC Ile														2448
		CTA Leu														2496
GAT Asp																2544

											AAA Lys				2592
											GAT Asp				2640
											GAT Asp			AGT Ser	2688
											GAT Asp				2736
											AAC Asn 925				2784
											AAT Asn				2832
											AAT Asn				2880
											TTA Leu				2928
											AAG Lys				2976
							Gly				CTT Leu 1005	Tyr			3024
	Leu					Ser					GGT Gly)				3072
Ser					Phe					Cys	AGT Ser				3120
				Tyr					Asp		GAA Glu			Glu	3168
			Thr					Glu			ACA Thr		Ile		3216
		Trp					Leu				GAA Glu 1085	Tyr			3264

Leu	AAT Asn 109	Val	TTA Leu	AAA Lys	Pro	AAT Asn 109	Asn	TTT	ATT Ile	GAT Asp	AGG Arg 110	Arg	AAA Lys	GAT Asp	TCT Ser	3312
ACT Thr 110	Leu	AGC Ser	ATT Ile	AAT Asn	AAT Asn 111	Ile	AGA Arg	AGC Ser	ACT Thr	ATT Ile 111	Leu	TTA Leu	GCT Ala	AAT Asn	AGA Arg 1120	3360
TTA Leu	TAT Tyr	AGT Ser	GGA Gly	ATA Ile 1125	Lys	GTT Val	AAA Lys	ATA Ile	CAA Gln 113	Arg	GTT Val	AAT Asn	AAT Asn	AGT Ser 113	Ser	3408
ACT Thr	AAC Asn	GAT Asp	AAT Asn 1140	Leu	GTT Val	AGA Arg	AAG Lys	AAT Asn 114	Asp	CAG Gln	GTA Val	TAT Tyr	ATT Ile 1150	Asn	TTT Phe	3456
GTA Val	GCC Ala	AGC Ser 115	AAA Lys	ACT Thr	CAC His	TTA Leu	TTT Phe 1160	Pro	TTA Leu	TAT Tyr	GCT Ala	GAT Asp 116	Thr	GCT Ala	ACC Thr	3504
ACA Thr	AAT Asn 1170	Lys	GAG Glu	AAA Lys	ACA Thr	ATA Ile 1175	Lys	ATA Ile	TCA Ser	TCA Ser	TCT Ser 118	Gly	AAT Asn	AGA Arg	TTT Phe	3552
AAT Asn 1185	Gln	GTA Val	GTA Val	GTT Val	ATG Met 1190	Asn	TCA Ser	GTA Val	GGA Gly	AAT Asn 1195	Asn	TGT Cys	ACA Thr	ATG Met	AÁT Asn 1200	3600
TTT Phe	AAA Lys	AAT Asn	AAT Asn	AAT Asn 1205	Gly	AAT Asn	AAT Asn	ATT Ile	GGG Gly 1210	Leu.	TTA Leu	GGT Gly	TTC Phe	AAG Lys 1215	Ala	3648
GAT Asp	ACT Thr	GTA Val	GTT Val 1220	Ala	AGT Ser	ACT Thr	TGG Trp	TAT Tyr 1225	Tyr	ACA Thr	CAT His	ATG Met	AGA Arg 1230	Asp	CAT His	3696
ACA Thr	AAC Asn	AGC Ser 1235	AAT Asn	GGA Gly	TGT Cys	Phe	TGG Trp 1240	Asn	TTT Phe	ATT Ile	TCT Ser	GAA Glu 1249	Glu	CAT His	GGA Gly	3744
Trp		Glu	AAA Lys	TAA				:							٠	3759

- (2) INFORMATION FOR SEQ ID NO:52:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1252 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:52:

Met Pro Lys Ile Asn Ser Phe Asn Tyr Asn Asp Pro Val Asn Asp Arg

1 10 15

Thr Ile Leu Tyr Ile Lys Pro Gly Gly Cys Gln Glu Phe Tyr Lys Ser 20 25 30

Phe Asn Ile Met Lys Asn Ile Trp Ile Ile Pro Glu Arg Asn Val Ile 35 40 45

Gly Thr Thr Pro Gln Asp Phe His Pro Pro Thr Ser Leu Lys Asn Gly 50 55 60

Asp Ser Ser Tyr Tyr Asp Pro Asn Tyr Leu Gln Ser Asp Glu Glu Lys
65 70 75 80

Asp Arg Phe Leu Lys Ile Val Thr Lys Ile Phe Asn Arg Ile Asn Asn 85 90 95

Asn Leu Ser Gly Gly Ile Leu Leu Glu Glu Leu Ser Lys Ala Asn Pro 100 105 110

Tyr Leu Gly Asn Asp Asn Thr Pro Asp Asn Gln Phe His Ile Gly Asp 115 120 125

Ala Ser Ala Val Glu Ile Lys Phe Ser Asn Gly Ser Gln Asp Ile Leu 130 135 140

Leu Pro Asn Val Ile Ile Met Gly Ala Glu Pro Asp Leu Phe Glu Thr 145 150 155 160

Asn Ser Ser Asn Ile Ser Leu Arg Asn Asn Tyr Met Pro Ser Asn His 165 170 175

Gly Phe Gly Ser Ile Ala Ile Val Thr Phe Ser Pro Glu Tyr Ser Phe 180 185 190

Arg Phe Asn Asp Asn Ser Met Asn Glu Phe Ile Gln Asp Pro Ala Leu 195 200 205

Thr Leu Met His Glu Leu Ile His Ser Leu His Gly Leu Tyr Gly Ala 210 215 220

Lys Gly Ile Thr Thr Lys Tyr Thr Ile Thr Gln Lys Gln Asn Pro Leu 225 230 235 240

Ile Thr Asn Ile Arg Gly Thr Asn Ile Glu Glu Phe Leu Thr Phe Gly
245 250 255

Gly Thr Asp Leu Asn Ile Ile Thr Ser Ala Gln Ser Asn Asp Ile Tyr 260 265 270

Thr Asn Leu Leu Ala Asp Tyr Lys Lys Ile Ala Ser Lys Leu Ser Lys 275 280 285

Val Gln Val Ser Asn Pro Leu Leu Asn Pro Tyr Lys Asp Val Phe Glu 290 295 300

Ala Lys Tyr Gly Leu Asp Lys Asp Ala Ser Gly Ile Tyr Ser Val Asn 305 310 315 320

Ile Asn Lys Phe Asn Asp Ile Phe Lys Lys Leu Tyr Ser Phe Thr Glu 325 330 335

Phe Asp Leu Ala Thr Lys Phe Gln Val Lys Cys Arg Gln Thr Tyr Ile 340 345 350

Gly Gln Tyr Lys Tyr Phe Lys Leu Ser Asn Leu Leu Asn Asp Ser Ile 355 360 355

Tyr Asn Ile Ser Glu Gly Tyr Asn Ile Asn Asn Leu Lys Val Asn Phe Arg Gly Gln Asn Ala Asn Leu Asn Pro Arg Ile Ile Thr Pro Ile Thr 390 Gly Arg Gly Leu Val Lys Lys Ile Ile Arg Phe Cys Lys Asn Ile Val Ser Val Lys Gly Ile Arg Lys Ser Ile Cys Ile Glu Ile Asn Asn Gly Glu Leu Phe Phe Val Ala Ser Glu Asn Ser Tyr Asn Asp Asp Asn Ile Asn Thr Pro Lys Glu Ile Asp Asp Thr Val Thr Ser Asn Asn Asn Tyr 455 Glu Asn Asp Leu Asp Gln Val Ile Leu Asn Phe Asn Ser Glu Ser Ala 470 Pro Gly Leu Ser Asp Glu Lys Leu Asn Leu Thr Ile Gln Asn Asp Ala 490 Tyr Ile Pro Lys Tyr Asp Ser Asn Gly Thr Ser Asp Ile Glu Gln His Asp Val Asn Glu Leu Asn Val Phe Phe Tyr Leu Asp Ala Gln Lys Val Pro Glu Gly Glu Asn Asn Val Asn Leu Thr Ser Ser Ile Asp Thr Ala 535 Leu Leu Glu Gln Pro Lys Ile Tyr Thr Phe Phe Ser Ser Glu Phe Ile Asn Asn Val Asn Lys Pro Val Gln Ala Ala Leu Phe Val Ser Trp Ile 570 Gln Gln Val Leu Val Asp Phe Thr Thr Glu Ala Asn Gln Lys Ser Thr 580 585 Val Asp Lys Ile Ala Asp Ile Ser Ile Val Val Pro Tyr Ile Gly Leu 600 Ala Leu Asn Ile Gly Asn Glu Ala Gln Lys Gly Asn Phe Lys Asp Ala Leu Glu Leu Leu Gly Ala Gly Ile Leu Leu Glu Phe Glu Pro Glu Leu 630 635 Leu Ile Pro Thr Ile Leu Val Phe Thr Ile Lys Ser Phe Leu Gly Ser Ser Asp Asn Lys Asn Lys Val Ile Lys Ala Ile Asn Asn Ala Leu Lys Glu Arg Asp Glu Lys Trp Lys Glu Val Tyr Ser Phe Ile Val Ser Asn 680 Trp Met Thr Lys Ile Asn Thr Gln Phe Asn Lys Arg Lys Glu Gln Met

Tyr Gln Ala Leu Gln Asn Gln Val Asn Ala Ile Lys Thr Ile Ile Glu 705 710 715 720

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- Ser Lys Tyr Asn Ser Tyr Thr Leu Glu Glu Lys Asn Glu Leu Thr Asn 725 730 735
- Lys Tyr Asp Ile Lys Gln Ile Glu Asn Glu Leu Asn Gln Lys Val Ser 740 745 750
- Ile Ala Met Asn Asn Ile Asp Arg Phe Leu Thr Glu Ser Ser Ile Ser 755 760 765
- Tyr Leu Met Lys Leu Ile Asn Glu Val Lys Ile Asn Lys Leu Arg Glu 770 780
- Tyr Asp Glu Asn Val Lys Thr Tyr Leu Leu Asn Tyr Ile Ile Gln His 785 790 795 800
- Gly Ser Ile Leu Gly Glu Ser Gln Gln Glu Leu Asn Ser Met Val Thr 805 810 815
- Asp Thr Leu Asn Asn Ser Ile Pro Phe Lys Leu Ser Ser Tyr Thr Asp 820 825 830
- Asp Lys Ile Leu Ile Ser Tyr Phe Asn Lys Phe Phe Lys Arg Ile Lys 835 840 845
- Ser Ser Ser Val Leu Asn Met Arg Tyr Lys Asn Asp Lys Tyr Val Asp 850 860
- Thr Ser Gly Tyr Asp Ser Asn Ile Asn Ile Asn Gly Asp Val Tyr Lys 865 870 875 880
- Tyr Pro Thr Asn Lys Asn Gln Phe Gly Ile Tyr Asn Asp Lys Leu Ser 885 890 895
- Glu Val Asn Ile Ser Gln Asn Asp Tyr Ile Ile Tyr Asp Asn Lys Tyr 900 905 910
- Lys Asn Phe Ser Ile Ser Phe Trp Val Arg Ile Pro Asn Tyr Asp Asn 915 920 925
- Lys Ile Val Asn Val Asn Asn Glu Tyr Thr Ile Ile Asn Cys Met Arg 930 935 940
- Asp Asn Asn Ser Gly Trp Lys Val Ser Leu Asn His Asn Glu Ile Ile 945 950 955 960
- Trp Thr Leu Gln Asp Asn Ala Gly Ile Asn Gln Lys Leu Ala Phe Asn 965 970 975
- Tyr Gly Asn Ala Asn Gly Ile Ser Asp Tyr Ile Asn Lys Trp Ile Phe 980 985 990
- Val Thr Ile Thr Asn Asp Arg Leu Gly Asp Ser Lys Leu Tyr Ile Asn 995 1000 1005
- Gly Asn Leu Ile Asp Gln Lys Ser Ile Leu Asn Leu Gly Asn Ile His 1010 1015 1020
- Val Ser Asp Asn Ile Leu Phe Lys Ile Val Asn Cys Ser Tyr Thr Arg 1025 1030 1035 1040

Tyr Ile Gly Ile Arg Tyr Phe Asn Ile Phe Asp Lys Glu Leu Asp Glu 1045 1050 1055

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Thr Glu Ile Gln Thr Leu Tyr Ser Asn Glu Pro Asn Thr Asn Ile Leu 1060 1065 1070

Lys Asp Phe Trp Gly Asn Tyr Leu Leu Tyr Asp Lys Glu Tyr Tyr Leu 1075 1080 1085

Leu Asn Val Leu Lys Pro Asn Asn Phe Ile Asp Arg Arg Lys Asp Ser 1090 1095 1100

Thr Leu Ser Ile Asn Asn Ile Arg Ser Thr Ile Leu Leu Ala Asn Arg 1105 1110 1115 1120

Leu Tyr Ser Gly Ile Lys Val Lys Ile Gln Arg Val Asn Asn Ser Ser 1125 1130 1135

Thr Asn Asp Asn Leu Val Arg Lys Asn Asp Gln Val Tyr Ile Asn Phe
1140 1145 1150

Val Ala Ser Lys Thr His Leu Phe Pro Leu Tyr Ala Asp Thr Ala Thr 1155 1160 1165

Thr Asn Lys Glu Lys Thr Ile Lys Ile Ser Ser Ser Gly Asn Arg Phe 1170 1175 1180

Asn Gln Val Val Wet Asn Ser Val Gly Asn Asn Cys Thr Met Asn 1185 1190 1195 1200

Phe Lys Asn Asn Gly Asn Asn Ile Gly Leu Leu Gly Phe Lys Ala 1205 1210 1215

Asp Thr Val Val Ala Ser Thr Trp Tyr Tyr Thr His Met Arg Asp His 1220 1225 1230

Thr Asn Ser Asn Gly Cys Phe Trp Asn Phe Ile Ser Glu Glu His Gly 1235 1240 1245

Trp Gln Glu Lys 1250

- (2) INFORMATION FOR SEQ ID NO:53:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1463 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid
 (A) DESCRIPTION: /desc = "DNA"
 - (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 108..1460

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:53:

AGATCTCGAT CCCG	CGAAAT TAATACGAC	T CACTATAGGG	GAATTGTGAG CGGA	TAACAA 60
TTCCCCTCTA GAAA	TAATTT TGTTTAACT	T TAAGAAGGAG	ATATACC ATG GGC Met Gly 1	
CAT CAT CAT CAT His His His His 5	CAT CAT CAT CAT His His His His 10	CAC AGC AGC His Ser Ser	GGC CAT ATC GAA Gly His Ile Glu 15	GGT 164 Gly
	AGC ATG GCT CTT Ser Met Ala Leu 25			
	TTT AAT AAG TTC Phe Asn Lys Phe 40			
GTT TTA AAT ATG Val Leu Asn Met 55	AGA TAT AAA AAT Arg Tyr Lys Asn	GAT AAA TAC Asp Lys Tyr 60	GTA GAT ACT TCA Val Asp Thr Ser 65	GGA 308 Gly
TAT GAT TCA AAT Tyr Asp Ser Asn 70	ATA AAT ATT AAT Ile Asn Ile Asn 75	GGA GAT GTA Gly Asp Val	TAT AAA TAT CCA Tyr Lys Tyr Pro 80	ACT 356 Thr
AAT AAA AAT CAA Asn Lys Asn Gln 85	TTT GGA ATA TAT Phe Gly Ile Tyr 90	AAT GAT AAA Asn Asp Lys	CTT AGT GAA GTT Leu Ser Glu Val 95	AAT 404 Asn
ATA TCT CAA AAT Ile Ser Gln Asn 100	GAT TAC ATT ATA Asp Tyr Ile Ile 105	TAT GAT AAT Tyr Asp Asn 110	AAA TAT AAA AAT Lys Tyr Lys Asn	TTT 452 Phe 115
AGT ATT AGT TTT Ser Ile Ser Phe	TGG GTA AGA ATT Trp Val Arg Ile 120	CCT AAC TAT Pro Asn Tyr 125	GAT AAT AAG ATA Asp Asn Lys Ile 130	GTA 500 Val
AAT GTT AAT AAT Asn Val Asn Asn 135	GAA TAC ACT ATA Glu Tyr Thr Ile	ATA AAT TGT Ile Asn Cys 140	ATG AGG GAT AAT Met Arg Asp Asn 145	AAT 548 Asn
TCA GGA TGG AAA Ser Gly Trp Lys 150	GTA TCT CTT AAT Val Ser Leu Asn 155	His Asn Glu	ATA ATT TGG ACA Ile Ile Trp Thr 160	TTG 596 Leu
CAA GAT AAT TCA Gln Asp Asn Ser 165	GGA ATT AAT CAA Gly Ile Asn Gln 170	AAA TTA GCA Lys Leu Ala	TTT AAC TAT GGT Phe Asn Tyr Gly 175	AAC 644 Asn
GCA AAT GGT ATT Ala Asn Gly Ile 180	TCT GAT TAT ATA Ser Asp Tyr Ile 185	AAT AAG TGG Asn Lys Trp 190	ATT TTT GTA ACT Ile Phe Val Thr	ATA 692 Ile 195
ACT AAT GAT AGA Thr Asn Asp Arg	TTA GGA GAT TCT Leu Gly Asp Ser 200	AAA CTT TAT Lys Leu Tyr 205	ATT AAT GGA AAT Ile Asn Gly Asn 210	Leu

ATA Ile	GAT Asp	AAA Lys	AAA Lys 215	TCA Ser	ATT Ile	TTA Leu	AAT Asn	TTA Leu 220	GGT Gly	AAT Asn	ATT Ile	CAT His	GTT Val 225	AGT Ser	GAC Asp	788
AAT Asn	ATA Ile	TTA Leu 230	TTT Phe	AAA Lys	ATA Ile	GTT Val	AAT Asn 235	TGT Cys	AGT Ser	TAT Tyr	ACA Thr	AGA Arg 240	TAT Tyr	ATT Ile	GGT Gly	836
ATT Ile	AGA Arg 245	TAT Tyr	TTT Phe	AAT Asn	ATT Ile	TTT Phe 250	GAT Asp	AAA Lys	GAA Glu	TTA Leu	GAT Asp 255	GAA Glu	ACA Thr	GAA Glu	ATT Ile	884
					AAT Asn 265											932
					CTT Leu											980
					TTT Phe											1028
					AGC Ser											1076
					ATA Ile											1124
					AAT Asn 345											1172
			Leu		CCA Pro											1220
GAG Glu																1268
GTA Val																1316
AAT Asn																1364
GCT Ala 420																1412
GGA Gly								Glu								1460
TAA																1463

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(2) INFORMATION FOR SEQ ID NO:54:

(i) SEQUENCE CHARACTERISTICS:

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- (A) LENGTH: 451 amino acids
- (B) TYPE: amino acid
- (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:54:
- Met Gly His His His His His His His His His Ser Ser Gly His

 1 10 15
- Ile Glu Gly Arg His Met Ala Ser Met Ala Leu Ser Ser Tyr Thr Asp
 20 25 30
- Asp Lys Ile Leu Ile Ser Tyr Phe Asn Lys Phe Phe Lys Arg Ile Lys
 35 40 45
- Ser Ser Ser Val Leu Asn Met Arg Tyr Lys Asn Asp Lys Tyr Val Asp 50 55 60
- Thr Ser Gly Tyr Asp Ser Asn Ile Asn Ile Asn Gly Asp Val Tyr Lys
 65 70 75 80
- Tyr Pro Thr Asn Lys Asn Gln Phe Gly Ile Tyr Asn Asp Lys Leu Ser 85 90 95
- Glu Val Asn Ile Ser Gln Asn Asp Tyr Ile Ile Tyr Asp Asn Lys Tyr 100 105 110
- Lys Asn Phe Ser Ile Ser Phe Trp Val Arg Ile Pro Asn Tyr Asp Asn 115 120 125
- Lys Ile Val Asn Val Asn Asn Glu Tyr Thr Ile Ile Asn Cys Met Arg 130 135 140
- Asp Asn Asn Ser Gly Trp Lys Val Ser Leu Asn His Asn Glu Ile Ile 145 150 155 160
- Trp Thr Leu Gln Asp Asn Ser Gly Ile Asn Gln Lys Leu Ala Phe Asn 170 175
- Tyr Gly Asn Ala Asn Gly Ile Ser Asp Tyr Ile Asn Lys Trp Ile Phe 180 185 190
- Val Thr Ile Thr Asn Asp Arg Leu Gly Asp Ser Lys Leu Tyr Ile Asn 195 200 205
- Gly Asn Leu Ile Asp Lys Lys Ser Ile Leu Asn Leu Gly Asn Ile His 210 215 220
- Val Ser Asp Asn Ile Leu Phe Lys Ile Val Asn Cys Ser Tyr Thr Arg 225 230 235 240
- Tyr Ile Gly Ile Arg Tyr Phe Asn Ile Phe Asp Lys Glu Leu Asp Glu 245 250 255
- Thr Glu Ile Gln Thr Leu Tyr Asn Asn Glu Pro Asn Ala Asn Ile Leu 260 265 270

Lys	qeA	Phe 275	Trp	Gly	Asn	Tyr	Leu 280	Leu	Tyr	Asp	Lys	Glu 285	Tyr	Tyr	Leu	
Leu	Asn 290	Val	Leu	Lys	Pro	Asn 295	Asn	Phe	Ile	Asn	Arg 300	Arg	Thr	Asp	Ser	
Thr 305	Leu	Ser	Ile	Asn	Asn 310	Ile	Arg	Ser	Thr	Ile 315	Leu	Leu	Ala	Asn	Arg 320	
Leu	Tyr	Ser	Gly	Ile 325	Lys	Val	Lys	Ile	Gln 330	Arg	Val	Asn	Asn	Ser 335	Ser	
Thr	Asn	Asp	Asn 340	Leu	Val	Arg	Lys	Asn 345	Asp	Gln	Val	Tyr	Ile 350	Asn	Phe	
Val	Ala	Ser 355	Lys	Thr	His	Leu	Leu 360	Pro	Leu	Tyr	Ala	Asp 365	Thr	Ala	Thr	
Thr	Asn 370	Lys	Glu	Lys	Thr	Ile 375	Lys	Ile	Ser	Ser	Ser 380	Gly	Asn	Arg	Phe	
Asn 385	Gln	Val	Val	Val	Met 390	Asn	Ser	Val	Gly	Asn 395	Cys	Thr	Met	Asn	Phe 400	
Lys	Asn	Asn	Asn	Gly 405	Asn	Asn	Ile	Gly	Leu 410	Leu	Gly	Phe	Lys	Ala 415		
Thr	Val	Val	Ala 420	Ser	Thr	Trp	Tyr	Tyr 425	Thr	His	Met	Arg	Asp 430	Asn	Thr	
Asn	Ser	Asn 435	Gly	Phe	Phe	Trp	Asn 440	Phe	Ile	Ser	Glu	Glu 445	His	Gly	Trp	
Gln	Glu 450	Lys														
(2)	INFO	RMAT	rion	FOR	SEQ	ID N	10 : 5	5:								
	(i)	(I (C	A) LI B) TY C) ST	engti (PE : (Rani	HARAC H: 14 nucl DEDNE DGY:	72 k eic SS:	ase acid	pai:	:s						,	
	(ii)				(PE:											
	(ix)		A) N	AME/E	KEY:		146	53								
	(xi)	SE	QUEN	CE DE	ESCRI	PTIC	N: 5	SEQ :	D NO):55	:					
AGAT	CTC	GAT (CCG	CGAA	AT TA	ATAC	CGACT	CAC	TAT	AGGG	GAA?	rtgt	GAG (CGGA:	TAACAA	60
TTC	CCT	CTA (gaaa'	TAAT	TT TO	GTTT/	AACT	r tal	AGAA	GGAG	ATA:	TACC		GGC Gly		116
CAT His	CAT His 5	CAT His	CAT His	CAT His	CAT His	CAT His	CAT His	CAC His	AGC Ser	AGC Ser	GGC Gly 15	CAT His	ATC Ile	GAA Glu	GGT Gly	164

CGT Arg 20	CAT His	ATG Met	GCT Ala	AGC Ser	ATG Met 25	GCT Ala	CTT Leu	TÇT Ser	TCT Ser	TAT Tyr 30	ACA Thr	GAT Asp	GAT Asp	AAA Lys	ATT Ile 35		212
TTA Leu	ATT Ile	TCA Ser	TAT Tyr	TTT Phe 40	AAT Asn	AAA Lys	TTC Phe	TTT Phe	AAG Lys 45	AGA Arg	ATT Ile	AAA Lys	AGT Ser	AGT Ser 50	TCA Ser		260
															GGA Gly		308
					AAT Asn												356
					GGA Gly												404
					TAC Tyr 105												452
					GTA Val			Pro									500
					TAC Tyr												548
					TCT Ser												596
					ATT Ile											·	644
					GAT Asp 185												692
					GGA Gly												740
					ATT Ile										GAC Asp		788
					ATA Ile												836
					ATT Ile												884

(

						GAA Glu										932
						TAT Tyr										980
						ATT Ile										1028
ATT Ile	AAT Asn	AAT Asn 310	ATA Ile	AGA Arg	AGC Ser	ACT Thr	ATT Ile 315	CTT Leu	TTA Leu	GCT Ala	AAT Asn	AGA Arg 320	TTA Leu	TAT Tyr	AGT Ser	1076
GGA Gly	ATA Ile 325	AAA Lys	GTT Val	AAA Lys	ATA Ile	CAA Gln 330	AGA Arg	GTT Val	AAT Asn	AAT Asn	AGT Ser 335	AGT Ser	ACT Thr	AAC Asn	GAT Asp	1124
						GAT Asp										1172
						TTA Leu										1220
						TCA Ser										1268
						GGA Gly										1316
						GGG Gly 410										1364
						TAT Tyr										1412
						TTT Phe										1460
AAA Lys	TAAA	AGCI	T													1472

(2) INFORMATION FOR SEQ ID NO:56:

(i) SEQUENCE CHARACTERISTICS:

(A) LENGTH: 452 amino acids

(B) TYPE: amino acid

(D) TOPOLOGY: linear

(ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:56:

Met Gly His His His His His His His His His Ser Ser Gly His

1 10 15

Ile Glu Gly Arg His Met Ala Ser Met Ala Leu Ser Ser Tyr Thr Asp
20 25 30

Asp Lys Ile Leu Ile Ser Tyr Phe Asn Lys Phe Phe Lys Arg Ile Lys 35 40 45

Ser Ser Ser Val Leu Asn Met Arg Tyr Lys Asn Asp Lys Tyr Val Asp 50 55 60

Thr Ser Gly Tyr Asp Ser Asn Ile Asn Ile Asn Gly Asp Val Tyr Lys
65 70 75 80

Tyr Pro Thr Asn Lys Asn Gln Phe Gly Ile Tyr Asn Asp Lys Leu Ser 85 90 95

Glu Val Asn Ile Ser Gln Asn Asp Tyr Ile Ile Tyr Asp Asn Lys Tyr
100 105 110

Lys Asn Phe Ser Ile Ser Phe Trp Val Arg Ile Pro Asn Tyr Asp Asn 115 120 125

Lys Ile Val Asn Val Asn Asn Glu Tyr Thr Ile Ile Asn Cys Met Arg

Asp Asn Asn Ser Gly Trp Lys Val Ser Leu Asn His Asn Glu Ile Ile 145 150 155 160

Trp Thr Leu Gln Asp Asn Ala Gly Ile Asn Gln Lys Leu Ala Phe Asn 165 170 175

Tyr Gly Asn Ala Asn Gly Ile Ser Asp Tyr Ile Asn Lys Trp Ile Phe 180 185 190

Val Thr Ile Thr Asn Asp Arg Leu Gly Asp Ser Lys Leu Tyr Ile Asn 195 200 205

Gly Asn Leu Ile Asp Gln Lys Ser Ile Leu Asn Leu Gly Asn Ile His 210 215 220

Val Ser Asp Asn Ile Leu Phe Lys Ile Val Asn Cys Ser Tyr Thr Arg 225 230 235 240

Tyr Ile Gly Ile Arg Tyr Phe Asn Ile Phe Asp Lys Glu Leu Asp Glu 245 250 255

Thr Glu Ile Gln Thr Leu Tyr Ser Asn Glu Pro Asn Thr Asn Ile Leu 260 265 270

Lys Asp Phe Trp Gly Asn Tyr Leu Leu Tyr Asp Lys Glu Tyr Tyr Leu 275 280 285

Leu Asn Val Leu Lys Pro Asn Asn Phe Ile Asp Arg Arg Lys Asp Ser 290 295 300

Thr Leu Ser Ile Asn Asn Ile Arg Ser Thr Ile Leu Leu Ala Asn Arg 305 310 315 320

Leu	Tyr	Ser	Gly	Ile 325	Lys	Val	Lys	Ile	Gln 330	Arg	Val	Asn	Asn	Ser 335	Ser	
Thr	Asn	Asp	Asn 340	Leu	Val	Arg	Lys	Asn 345	Asp	Gln	Val	Tyr	Ile 350	Asn	Phe	
Val	Ala	Ser 355	Lys	Thr	His	Leu	Phe 360	Pro	Leu	Tyr	Ala	Asp 365	Thr	Ala	Thr	
Thr	Asn 370	Lys	Glu	Lys	Thr	Ile 375	Lys	Ile	Ser	Ser	Ser 380	Gly	Asn	Arg	Phe	
Asn 385	Gln	Val	Val	Val	Met 390	Asn	Ser	Val	Gly	Asn 395	Asn	Cys	Thr	Met	Asn 400	
Phe	Lys	Asn	Asn	Asn 405	Gly	Asn	Asn	Ile	Gly 410		Leu	Gly	Phe	Lys 415	Ala	
Asp	Thr	Val	Val 420	Ala	Ser	Thr	Trp	Tyr 425	Tyr	Thr	His	Met	Arg 430	Asp	His	
Thr	Asn	Ser 435	Asn	Gly	Cys	Phe	Trp 440	Asn	Phe	Ile	Ser	Glu 445	Glu	His	Gly	
Trp	Gln 450	Glu	Lys													
(2)	INFO	RMAI	CION	FOR	SEQ	ID N	ro : 57	':								
	(i)	(E (C	() LE 3) TY C) ST	ENGTI (PE : [RANI	I: 31 nucl EDNE	TERI bas eic ESS: line	e pa acid	irs l							•	
	(ii)								c ac							
	(xi)	SEC	UENC	E DE	ESCRI	PTIC	N: S	EQ 1	D NO	57:						
CGCC	CATGO	CT C	CTTTC	CTTCT	TA T	CACAC	ATG	Т								31
(2)	INFO	RMAT	CION	FOR	SEQ	ID N	rO : 58	B:								
	(i)	() () ()	A) LE 3) TY C) SI	CE CH ENGTH (PE: TRANI	i: 29 nucl EDNE	TERI bas eic SS: line	e pa acid sing	lirs								
	(ii)	MOI ()	LECUI	ESCRI	(PE:	othe	r nu desc	clei	ic ac	id'						
	(xi)	SEC	QUENC	E DE	ESCRI	PTIC	N: S	EQ 1	D NO):58:						

GCAAGCTTTT ATTTTTCTTG CCATCCATG

29

	(i	((A) L (B) T (C) S	ENGT YPE : TRAN	H: 3 nuc DEDN	CTER 876 leic ESS:	base aci dou	pai d	.rs							
	(ii) MO	LECU	LE T	YPE:	DNA	(ge	nomi	c)							
	(ix	(AME/		CDS										
	(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:59	:					
ATG Met 1	Pro	ATA Ile	ACA Thr	ATT Ile 5	AAC Asn	AAC Asn	TTT Phe	AAT Asn	TAT Tyr 10	TCA Ser	GAT Asp	CCT	GTT Val	GAT Asp 15	AAT Asn	4.8
AAA Lys	AAT Asn	ATT Ile	TTA Leu 20	TAT Tyr	TTA Leu	GAT Asp	ACT Thr	CAT His 25	TTA Leu	AAT Asn	ACA Thr	CTA Leu	GCT Ala 30	AAT Asn	GAG Glu	96
CCT Pro	GAA Glu	AAA Lys 35	GCC Ala	TTT Phe	CGC Arg	ATT Ile	ACA Thr 40	GGA Gly	AAT Asn	ATA Ile	TGG Trp	GTA Val 45	ATA Ile	CCT Pro	GAT Asp	144
AGA Arg	TTT Phe 50	TCA Ser	AGA Arg	AAT Asn	TCT Ser	AAT Asn 55	CCA Pro	AAT Asn	TTA Leu	AAT Asn	AAA Lys 60	CCT Pro	CCT Pro	CGA Arg	GTT Val	192
ACA Thr 65	AGC Ser	CCT Pro	AAA Lys	AGT Ser	GGT Gly 70	TAT	TAT Tyr	GAT Asp	CCT Pro	AAT Asn 75	TAT Tyr	TTG Leu	AGT Ser	ACT Thr	GAT Asp 80	240
TCT Ser	GAC Asp	AAA Lys	GAT Asp	ACA Thr 85	TTT Phe	TTA Leu	AAA Lys	GAA Glu	ATT Ile 90	ATA Ile	AAG Lys	TTA Leu	TTT Phe	AAA Lys 95	AGA Arg	288
ATT [le	AAT Asn	TCT Ser	AGA Arg 100	GAA Glu	ATA Ile	GGA Gly	GAA Glu	GAA Glu 105	TTA Leu	ATA Ile	TAT Tyr	AGA Arg	CTT Leu 110	TCG Ser	ACA Thr	336
						AAT Asn			Thr		Ile	Asn			GAT Asp	384
TTT he	GAT Asp 130	GTA Val	GAT Asp	TTT Phe	AAC Asn	AGT Ser 135	GTT Val	GAT Asp	GTT Val	AAA Lys	ACT Thr 140	AGA Arg	CAA Gln	GGT Gly	AAC Asn	432
						AGC Ser										480
						GAT Asp										528

(2) INFORMATION FOR SEQ ID NO:59:

AAC Asn	AAT Asn	ACT Thr	TTT Phe 180	GCG Ala	GCA Ala	CAA Gln	GAA Glu	GGA Gly 185	TTT	GGT	GCT Ala	TTA Leu	TCA Ser 190	Ile	ATT Ile	. 5	76
TCA Ser	ATA Ile	TCA Ser 195	CCT Pro	AGA Arg	TTT Phe	ATG Met	CTA Leu 200	Thr	TAT	AGT Ser	AAT Asn	GCA Ala 205	ACT Thr	AAT Asn	GAT Asp	6	24
GTA Val	GGA Gly 210	GAG Glu	GGT Gly	AGA Arg	TTT Phe	TCT Ser 215	AAG Lys	TCT Ser	GAA Glu	TTT Phe	TGC Cys 220	ATG Met	GAT Asp	CCA Pro	ATA Ile	6	72
CTA Leu 225	ATT Ile	TTA Leu	ATG Met	CAT His	GAA Glu 230	CTT Leu	AAT Asn	CAT His	GCA Ala	ATG Met 235	CAT His	AAT Asn	TTA Leu	TAT Tyr	GGA Gly 240	7:	20
ATA Ile	GCT Ala	ATA Ile	CCA Pro	AAT Asn 245	GAT Asp	CAA Gln	ACA Thr	ATT Ile	TCA Ser 250	TCT Ser	GTA Val	ACT Thr	AGT Ser	AAT Asn 255	Ile	76	68
TTT Phe	TAT Tyr	TCT Ser	CAA Gln 260	TAT Tyr	AAT Asn	GTG Val	AAA Lys	TTA Leu 265	GAG Glu	TAT Tyr	GCA Ala	GAA Glu	ATA Ile 270	TAT Tyr	GCA Ala	83	16
														AAA Lys		8 6	54
														AGA Arg		91	12
														ATA Ile		96	50
														GAA Glu 335		100	8
														TAT Tyr		105	56
GAA Glu	CTT Leu	ACA Thr 355	Gln	ATA Ile	TTT Phe	ACA Thr	GAA Glu 360	TTT Phe	AAC Asn	TAC Tyr	GCT Ala	AAA Lys 365	ATA Ile	TAT Tyr	AAT Asn	110	14
														GTT Val		115	52
														TTT Phe		120	00
														TTA Leu 415		124	8

CGA Arg	AAT Asn	CCA Pro	GCA Ala 420	Leu	AGA Arg	AAA Lys	GTC Val	AAT Asn 425	Pro	GAA Glu	AAT Asn	ATG Met	CTT Leu 430	TAT Tyr	TTA Leu	1296
TTT Phe	ACA Thr	AAA Lys 435	Phe	TGT	CAT	AAA Lys	GCA Ala 440	Ile	GAT Asp	GGT Gly	AGA Arg	TCA Ser 445	TTA Leu	TAT Tyr	AAT Asn	1344
AAA Lys	ACA Thr 450	TTA Leu	GAT Asp	TGT Cys	AGA Arg	GAG Glu 455	CTT Leu	TTA Leu	GTT Val	AAA Lys	AAT Asn 460	ACT Thr	GAC Asp	TTA Leu	CCC Pro	1392
TTT Phe 465	Ile	GGT Gly	GAT Asp	ATT Ile	AGT Ser 470	GAT Asp	GTT Val	AAA Lys	ACT Thr	GAT Asp 475	ATA Ile	TTT Phe	TTA Leu	AGA Arg	AAA Lys 480	1440
GAT Asp	ATT Ile	AAT Asn	GAA Glu	GAA Glu 485	ACT Thr	GAA Glu	GTT Val	ATA Ile	TAC Tyr 490	TAT Tyr	CCG Pro	GAC Asp	TAA Asn	GTT Val 495	TCA Ser	1488
GTA Val	GAT Asp	CAA Gln	GTT Val 500	ATT Ile	CTC Leu	AGT Ser	AAG Lys	AAT Asn 505	ACC Thr	TCA Ser	GAA Glu	CAT His	GGA Gly 510	CAA Gln	CTA Leu	1536
		TTA Leu 515														1584
		CAA Gln														1632
AAT Asn 545	TCT Ser	TAT Tyr	TAT Tyr	TAC Tyr	CTA Leu 550	GAA Glu	TCT Ser	CAA Gln	AAA Lys	CTA Leu 555	AGT Ser	GAT Asp	AAT Asn	GTT Val	GAA Glu 560	1680
		ACT Thr														1728
		TAT Tyr														1776
	Gln	GGT Gly 595														1824
TTT Phe	ACT Thr 610	ACA Thr	AAT Asn	ATT Ile	CTA Leu	AGA Arg 615	AAA Lys	GAT Asp	ACA Thr	TTA Leu	GAT Asp 620	AAA Lys	ATA Ile	TCA Ser	GAT Asp	1872
		GCT Ala							Pro							1920
		AGA Arg														1968

ACT Thr	ATT	TTA Leu	TTA Leu 660	GAA Glu	GCA Ala	TTT Phe	CCT Pro	GAA Glu 665	Phe	ACA Thr	ATA Ile	CCT Pro	GCA Ala 670	Leu	GGT	2016
GCA Ala	TTT	GTG Val 675	Ile	TAT Tyr	AGT Ser	AAG Lys	GTT Val 680	Gln	GAA Glu	AGA Arg	AAC Asn	GAG Glu 685	Ile	ATT	AAA Lys	2064
ACT Thr	ATA Ile 690	Asp	AAT Asn	TGT Cys	TTA Leu	GAA Glu 695	CAA Gln	AGG Arg	ATT	AAG Lys	AGA Arg 700	Trp	AAA Lys	GAT Asp	TCA Ser	2112
TAT Tyr 705	GAA Glu	TGG Trp	ATG Met	ATG Met	GGA Gly 710	ACG Thr	TGG Trp	TTA Leu	TCC Ser	AGG Arg 715	ATT Ile	ATT Ile	ACT	CAA Gln	TTT Phe 720	2160
AAT Asn	AAT Asn	ATA Ile	AGT Ser	TAT Tyr 725	CAA Gln	ATG Met	TAT Tyr	GAT Asp	TCT Ser 730	TTA Leu	AAT Asn	TAT Tyr	CAG Gln	GCA Ala 735	GGT	2208
GCA Ala	ATC Ile	AAA Lys	GCT Ala 740	AAA Lys	ATA Ile	GAT Asp	TTA Leu	GAA Glu 745	TAT Tyr	AAA Lys	AAA Lys	TAT	TCA Ser 750	GGA Gly	AGT Ser	2256
GAT Asp	AAA Lys	GAA Glu 755	AAT Asn	ATA Ile	AAA Lys	AGT Ser	CAA Gln 760	GTT Val	GAA Glu	AAT Asn	TTA Leu	AAA Lys 765	AAT Asn	AGT Ser	TTA Leu	2304
GAT Asp	GTA Val 770	AAA Lys	ATT Ile	TCG Ser	GAA Glu	GCA Ala 775	ATG Met	AAT Asn	AAT Asn	ATA Ile	AAT Asn 780	AAA Lys	TTT Phe	ATA Ile	CGA Arg	2352
GAA Glu 785	TGT Cys	TCC Ser	GTA Val	ACA Thr	TAT Tyr 790	TTA Leu	TTT Phe	AAA Lys	AAT Asn	ATG Met 795	TTA Leu	CCT Pro	AAA Lys	GTA Val	ATT Ile 800	2400
GAT Asp	GAA Glu	TTA Leu	AAT Asn	GAG Glu 805	TTT Phe	GAT Asp	CGA Arg	AAT Asn	ACT Thr 810	AAA Lys	GCA Ala	AAA Lys	TTA Leu	ATT Ile 815	AAT. Asn	2448
CTT Leu	ATA Ile	GAT Asp	AGT Ser 820	CAT His	AAT Asn	ATT Ile	ATT Ile	CTA Leu 825	GTT Val	GGT Gly	GAA Glu	GTA Val	GAT Asp 830	AAA Lys	TTA Leu	2496
AAA Lys	GCA Ala	AAA Lys 835	GTA Val	AAT Asn	AAT Asn	AGC Ser	TTT Phe 840	CAA Gln	AAT Asn	ACA Thr	ATA Ile	CCC Pro 845	TTT Phe	AAT Asn	ATT Ile	2544
												ATT Ile				2592
TTC Phe 865	AAT Asn	AAT Asn	ATT Ile	AAT Asn	GAT Asp 870	TCA Ser	AAA Lys	ATT Ile	TTG Leu	AGC Ser 875	CTA Leu	CAA Gln	AAC Asn	AGA Arg	AAA Lys 880	2640
			Val					Tyr				GTG Val				2688

GGC Gly	GAT Asp	GTT Val	Gln 900	Leu	AAT Asn	CCA Pro	ATA	TTI Phe 905		TTT	GAC Asp	TTT Phe	AAA Lys 910	TTA Leu	GGT	2736
AGT Ser	TCA Ser	GGG Gly 915	Glu	GAT Asp	AGA Arg	GGT Gly	AAA Lys 920	Val	ATA Ile	GTA Val	ACC Thr	CAG Gln 925	AAT Asn	GAA Glu	AAT Asn	2784
ATT Ile	GTA Val 930	TAT	AAT Asn	TCT Ser	ATG Met	TAT Tyr 935	GAA Glu	AGT Ser	TTT	AGC Ser	ATT Ile 940	AGT Ser	TTT Phe	TGG Trp	ATT Ile	2832
AGA Arg 945	ATA Ile	AAT Asn	AAA Lys	TGG	GTA Val 950	AGT Ser	AAT Asn	TTA Leu	CCT	GGA Gly 955	TAT Tyr	ACT Thr	ATA Ile	ATT Ile	GAT Asp 960	2880
AGT Ser	GTT Val	AAA Lys	AAT Asn	AAC Asn 965	TCA Ser	GGT Gly	TGG Trp	AGT Ser	ATA Ile 970	GGT Gly	ATT Ile	ATT Ile	AGT Ser	AAT Asn 975	TTT Phe	2928
TTA Leu	GTA Val	TTT Phe	ACT Thr 980	TTA Leu	AAA Lys	CAA Gln	AAT Asn	GAA Glu 985	GAT Asp	AGT Ser	GAA Glu	CAA Gln	AGT Ser 990	ATA Ile	AAT Asn	2976
TTT Phe	AGT Ser	TAT Tyr 995	GAT Asp	ATA Ile	TCA Ser	AAT Asn	AAT Asn 1000	Ala	CCT Pro	GGA Gly	TAC Tyr	AAT Asn 1005	Lys	TGG Trp	TTT Phe	3024
Phe	GTA Val 1010	Thr	GTT Val	ACT Thr	AAC Asn	AAT Asn 1019	Met	ATG Met	GGA Gly	AAT Asn	ATG Met 1020	Lys	ATT Ile	TAT Tyr	ATA Ile	3072
AAT Asn 1025	Gly	AAA Lys	TTA Leu	ATA Ile	GAT Asp 1030	Thr	ATA Ile	AAA Lys	GTT Val	AAA Lys 1035	Glu	CTA Leu	ACT Thr	GGA Gly	ATT Ile 1040	3120
AAT Asn	TTT Phe	AGC Ser	AAA Lys	ACT Thr 1045	Ile	ACA Thr	TTT Phe	GAA Glu	ATA Ile 1050	Asn	AÀA Lys	ATT Ile	CCA Pro	GAT Asp 1055	Thr	3168
GGT Gly	TTG Leu	ATT Ile	ACT Thr 1060	Ser	GAT Asp	TCT Ser	GAT Asp	AAC Asn 1069	ATC Ile	AAT Asn	ATG Met	TGG Trp	ATA Ile 1070	Arg	GAT Asp	3216
			Phe					Asp	GGT Gly				Asn			3264
Phe	AAT Asn 1090	Ser	TTG Leu	CAA Gln	TAT Tyr	ACT Thr 1095	Asn	GTT Val	GTA Val	AAA Lys	GAT Asp 1100	Tyr	TGG Trp	GGA Gly	AAT Asn	3312
GAT Asp 1105	Leu	AGA Arg	TAT Tyr	AAT Asn	AAA Lys 1110	Glu	TAT Tyr	TAT Tyr	ATG Met	GTT Val 1115	Asn	ATA Ile	GAT Asp	TAT Tyr	TTA Leu 1120	3360
					Ala				CAA Gln 1130	Ile			Asn		Arg	3408

AGA Arg	AAT Asn	AAT Asn	AAT Asn 114	Asp	TTC Phe	AAT Asn	GAA Glu	GGA Gly 114	Tyr	AAA Lys	ATT Ile	ATA Ile	ATA Ile 115	Lys	AGA Arg	3456
ATC Ile	AGA Arg	GGA Gly 115	Asn	ACA Thr	AAT Asn	GAT Asp	ACT Thr 1160	Arg	GTA Val	CGA Arg	GGA Gly	GGA Gly 116	Asp	ATT Ile	TTA Leu	3504
TAT Tyr	TTT Phe 1170	Asp	ATG Met	ACA Thr	ATT Ile	AAT Asn 1179	Asn	AAA Lys	GCA Ala	TAT Tyr	AAT Asn 1180	Leu	TTT Phe	ATG Met	AAG Lys	3552
AAT Asn 118	GAA Glu 5	ACT Thr	ATG Met	TAT Tyr	GCA Ala 1190	Asp	AAT Asn	CAT His	AGT Ser	ACT Thr 1199	Glu	GAT Asp	ATA Ile	TAT Tyr	GCT Ala 1200	3600
ATA Ile	GGT Gly	TTA Leu	AGA Arg	GAA Glu 1209	Gln	ACA Thr	AAG Lys	GAT Asp	ATA Ile 1210	Asn	GAT Asp	AAT Asn	ATT Ile	ATA Ile 1215	Phe	3648
CAA Gln	ATA Ile	CAA Gln	CCA Pro 1220	Met	AAT Asn	AAT Asn	ACT Thr	TAT Tyr 1225	Tyr	TAC Tyr	GCA Ala	TCT Sér	CAA Gln 1230	Ile	TTT Phe	3696
AAA Lys	TCA Ser	AAT Asn 1235	Phe	AAT Asn	GGA Gly	Glu	AAT Asn 1240	Ile	TCT Ser	GGA Gly	ATA Ile	TGT Cys 1245	Ser	ATA Ile	GGT Gly	3744
ACT Thr	TAT Tyr 1250	Arg	TTT Phe	AGA Arg	Leu	GGA Gly 1255	Gly	GAT Asp	TGG Trp	Tyr	AGA Arg 1260	His	AAT Asn	TAT Tyr	TTG Leu	3792
GTG Val 1265	CĊT Pro	ACT Thr	GTG Val	Lys	CAA Gln 1270	Gly	AAT Asn	TAT Tyr	GCT Ala	TCA Ser 1275	Leu	TTA. Leu	GAA Glu	Ser	ACA Thr 1280	3840
TCA Ser	ACT Thr	CAT His	Trp	GGT Gly 1285	Phe	GTA Val	CCT Pro	Val	AGT Ser 1290	Glu	TAA					3876
(2)	INFO	RMAT	ION	FOR	SEQ	ID N	0:60	:								

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1291 amino acids(B) TYPE: amino acid

 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:60:

Met Pro Ile Thr Ile Asn Asn Phe Asn Tyr Ser Asp Pro Val Asp Asn

Lys Asn Ile Leu Tyr Leu Asp Thr His Leu Asn Thr Leu Ala Asn Glu

Pro Glu Lys Ala Phe Arg Ile Thr Gly Asn Ile Trp Val Ile Pro Asp 35 40 45

Arg Phe Ser Arg Asn Ser Asn Pro Asn Leu Asn Lys Pro Pro Arg Val

65	261	PIO	БУЗ	261	70	Tyr	lyr	Asp	Pro	75	ıyı	rea	ser	Thr	Asp 80
Ser	Asp	Lys	Asp	Thr 85	Phe	Leu	Lys	Glu	Ile 90	Ile	Lys	Leu	Phe	Lys 95	Arg
Ile	Asn	Ser	Arg 100	Glu	Ile	Gly	Glu	Glu 105	Leu	Ile	Tyr	Arg	Leu 110	Ser	Thr
Asp	Ile	Pro 115	Phe	Pro	Gly	Asn	Asn 120	Asn	Thr	Pro	Ile	Asn 125	Thr	Phe	Asp
Phe	Asp 130	Val	Asp	Phe	Asn	Ser 135	Val	Asp	Val	Lys	Thr 140	Arg	Gln	Gly	Asn
Asn 145	Trp	Val	Lys	Thr	Gly 150	Ser	Ile	Asn	Pro	Ser 155	Val	Ile	Ile	Thr	Gly 160
Pro	Arg	Glu	Asn	Ile 165	Ile	Asp	Pro	Glu	Thr 170	Ser	Thr	Phe	Lys	Leu 175	Thr
Asn	Asn	Thr	Phe 180	Ala	Ala	Gln	Glu	Gly 185	Phe	Gly	Ala	Leu	Ser 190	Ile	Ile
Ser	Ile	Ser 195	Pro	Arg	Phe	Met	Leu 200	Thr	Tyr	Ser	Asn	Ala 205	Thr	Asn	Asp
Val	Gly 210	Glu	Gly	Arg	Phe	Ser 215	Lys	Ser	Glu	Phe	Cys 220	Met	Asp	Pro	Ile
Leu 225	Ile	Leu	Met	His	Glu 230	Leu	Asn	His	Ala	Met 235	His	Asn	Leu	Tyr	Gly 240
Ile	Ala	Ile	Pro	Asn 245	Asp	Gln	Thr	Ile	Ser 250	Ser	Val	Thr	Ser	Asn 255	Ile
Phe	Tyr	Ser	Gln 260	Tyr	Asn	Val	Lys	Leu 265	Glu	Tyr	Ala	Glu	Ile 270	Tyr	Ala
Phe	Gly	Gly 275	Pro	Thr	Ile	Asp	Leu 280	Ile	Pro	Lys	Ser	Ala 285		Lys	Tyr
Phe	Glu 290	Glu	Lys	Ala	Leu	Asp 295	Tyr	Tyr	Arg	Ser	Ile 300	Ala	Lys	Arg	Leu
Asn 305	Ser	Ile	Thr	Thr	Ala 310	Asn	Pro	Ser	Ser	Phe 315	Asn	Lys	Tyr	Ile	Gly 320
Glu	Tyr	Lys	Gln	Lys 325	Leu	Ile	Arg	Lys	Tyr 330	Arg	Phe	Val	Val	Glu 335	Ser
Ser	Gly	Glu	Val 340	Thr	Val	Asn	Arg	Asn 345	Lys	Phe	Val	Glu	Leu 350	Tyr	Asn
Glu	Leu	Thr 355	Gln	Ile	Phe	Thr	Glu 360	Phe	Asn	Tyr	Ala	Lys 365	Ile	Tyr	Asn
Val	Gln 370	Asn	Arg	Lys	Ile	Tyr 375	Leu	Ser	Asn	Val	Tyr 380	Thr	Pro	Val	Thr
Ala 385	Asn	Ile	Leu	Asp	Asp	Asn	Val	Tyr	Asp	Ile	Gln	Asn	Gly	Phe	Asn 400

Ile Pro Lys Ser Asn Leu Asn Val Leu Phe Met Gly Gln Asn Leu Ser Arg Asn Pro Ala Leu Arg Lys Val Asn Pro Glu Asn Met Leu Tyr Leu Phe Thr Lys Phe Cys His Lys Ala Ile Asp Gly Arg Ser Leu Tyr Asn Lys Thr Leu Asp Cys Arg Glu Leu Leu Val Lys Asn Thr Asp Leu Pro 455 Phe Ile Gly Asp Ile Ser Asp Val Lys Thr Asp Ile Phe Leu Arg Lys Asp Ile Asn Glu Glu Thr Glu Val Ile Tyr Tyr Pro Asp Asn Val Ser 490 Val Asp Gln Val Ile Leu Ser Lys Asn Thr Ser Glu His Gly Gln Leu Asp Leu Leu Tyr Pro Ser Ile Asp Ser Glu Ser Glu Ile Leu Pro Gly 520 Glu Asn Gln Val Phe Tyr Asp Asn Arg Thr Gln Asn Val Asp Tyr Leu Asn Ser Tyr Tyr Leu Glu Ser Gln Lys Leu Ser Asp Asn Val Glu Asp Phe Thr Phe Thr Arg Ser Ile Glu Glu Ala Leu Asp Asn Ser Ala Lys Val Tyr Thr Tyr Phe Pro Thr Leu Ala Asn Lys Val Asn Ala Gly 585 Val Gln Gly Gly Leu Phe Leu Met Trp Ala Asn Asp Val Val Glu Asp Phe Thr Thr Asn Ile Leu Arg Lys Asp Thr Leu Asp Lys Ile Ser Asp 615 Val Ser Ala Ile Ile Pro Tyr Ile Gly Pro Ala Leu Asn Ile Ser Asn Ser Val Arg Arg Gly Asn Phe Thr Glu Ala Phe Ala Val Thr Gly Val Thr Ile Leu Leu Glu Ala Phe Pro Glu Phe Thr Ile Pro Ala Leu Gly 665 Ala Phe Val Ile Tyr Ser Lys Val Gln Glu Arg Asn Glu Ile Ile Lys Thr Ile Asp Asn Cys Leu Glu Gln Arg Ile Lys Arg Trp Lys Asp Ser 695

Tyr Glu Trp Met Met Gly Thr Trp Leu Ser Arg Ile Ile Thr Gln Phe

Asn Asn Ile Ser Tyr Gln Met Tyr Asp Ser Leu Asn Tyr Gln Ala Gly

Ala Ile Lys Ala Lys Ile Asp Leu Glu Tyr Lys Lys Tyr Ser Gly Ser 740 745 750

Asp Lys Glu Asn Ile Lys Ser Gln Val Glu Asn Leu Lys Asn Ser Leu 755 760 765

Asp Val Lys Ile Ser Glu Ala Met Asn Asn Ile Asn Lys Phe Ile Arg 770 775 780

Glu Cys Ser Val Thr Tyr Leu Phe Lys Asn Met Leu Pro Lys Val Ile 785 790 795 800

Asp Glu Leu Asn Glu Phe Asp Arg Asn Thr Lys Ala Lys Leu Ile Asn 805 810 815

Leu Ile Asp Ser His Asn Ile Ile Leu Val Gly Glu Val Asp Lys Leu 820 825 830

Lys Ala Lys Val Asn Asn Ser Phe Gln Asn Thr Ile Pro Phe Asn Ile 835 840 845

Phe Ser Tyr Thr Asn Asn Ser Leu Leu Lys Asp Ile Ile Asn Glu Tyr 850 855

Phe Asn Asn Ile Asn Asp Ser Lys Ile Leu Ser Leu Gln Asn Arg Lys 865 870 875 880

Asn Thr Leu Val Asp Thr Ser Gly Tyr Asn Ala Glu Val Ser Glu Glu 885 890 895

Gly Asp Val Gln Leu Asn Pro Ile Phe Pro Phe Asp Phe Lys Leu Gly 900 905 910

Ser Ser Gly Glu Asp Arg Gly Lys Val Ile Val Thr Gln Asn Glu Asn 915 920 925

Ile Val Tyr Asn Ser Met Tyr Glu Ser Phe Ser Ile Ser Phe Trp Ile 930 935 940

Arg Ile Asn Lys Trp Val Ser Asn Leu Pro Gly Tyr Thr Ile Ile Asp 945 950 955 960

Ser Val Lys Asn Asn Ser Gly Trp Ser Ile Gly Ile Ile Ser Asn Phe 965 970 975

Leu Val Phe Thr Leu Lys Gln Asn Glu Asp Ser Glu Gln Ser Ile Asn 980 985 990

Phe Ser Tyr Asp Ile Ser Asn Asn Ala Pro Gly Tyr Asn Lys Trp Phe 995 1000 1005

Phe Val Thr Val Thr Asn Asn Met Met Gly Asn Met Lys Ile Tyr Ile 1010 1015 1020

Asn Gly Lys Leu Ile Asp Thr Ile Lys Val Lys Glu Leu Thr Gly Ile 1025 1030 1035 1040

Asn Phe Ser Lys Thr Ile Thr Phe Glu Ile Asn Lys Ile Pro Asp Thr 1045 1050 1055

Gly Leu Ile Thr Ser Asp Ser Asp Asn Ile Asn Met Trp Ile Arg Asp 1060 1065 1070 Phe Tyr Ile Phe Ala Lys Glu Leu Asp Gly Lys Asp Ile Asn Ile Leu 1075 1080 1085

Phe Asn Ser Leu Gln Tyr Thr Asn Val Val Lys Asp Tyr Trp Gly Asn 1090 1095 1100

Asp Leu Arg Tyr Asn Lys Glu Tyr Tyr Met Val Asn Ile Asp Tyr Leu 1105 1110 1115 1120

Asn Arg Tyr Met Tyr Ala Asn Ser Arg Gln Ile Val Phe Asn Thr Arg 1125 1130 1135

Arg Asn Asn Asn Asp Phe Asn Glu Gly Tyr Lys Ile Ile Ile Lys Arg 1140 1145 1150

Ile Arg Gly Asn Thr Asn Asp Thr Arg Val Arg Gly Gly Asp Ile Leu 1155 1160 1165

Tyr Phe Asp Met Thr Ile Asn Asn Lys Ala Tyr Asn Leu Phe Met Lys 1170 1175 1180

Asn Glu Thr Met Tyr Ala Asp Asn His Ser Thr Glu Asp Ile Tyr Ala 1185 1190 1195 1200

Ile Gly Leu Arg Glu Gln Thr Lys Asp Ile Asn Asp Asn Ile Ile Phe 1205 1210 1215

Gln Ile Gln Pro Met Asn Asn Thr Tyr Tyr Tyr Ala Ser Gln Ile Phe 1220 1225 1230

Lys Ser Asn Phe Asn Gly Glu Asn Ile Ser Gly Ile Cys Ser Ile Gly
1235 1240 1245

Thr Tyr Arg Phe Arg Leu Gly Gly Asp Trp Tyr Arg His Asn Tyr Leu 1250 1255 1260

Val Pro Thr Val Lys Gln Gly Asn Tyr Ala Ser Leu Leu Glu Ser Thr 1265 1270 1275 1280

Ser Thr His Trp Gly Phe Val Pro Val Ser Glu 1285 1290

(2) INFORMATION FOR SEQ ID NO:61:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1502 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 108..1493

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:61:

AGATCTCGAT CCCGCGAAA	AT TAATACGACT C	ACTATAGGG GAAT	TGTGAG CGGATAACAA 60
TTCCCCTCTA GAAATAATT	TT TGTTTAACTT TA	AAGAAGGAG ATATA	ACC ATG GGC CAT 116 Met Gly His 1
CAT CAT CAT CAT His His His His 5			
CGT CAT ATG GCT AGC Arg His Met Ala Ser 20			
TTC AAT AAT ATT AAT Phe Asn Asn Ile Asn 40			
AAT ACT TTA GTG GAT Asn Thr Leu Val Asp 55		c Asn Ala Glu \	
GGC GAT GTT CAG CTT Gly Asp Val Gln Leu 70			
AGT TCA GGG GAG GAT Ser Ser Gly Glu Asp 85			
ATT GTA TAT AAT TCT Ile Val Tyr Asn Ser 100			
AGA ATA AAT AAA TGG Arg Ile Asn Lys Trp 120			
AGT GTT AAA AAT AAC Ser Val Lys Asn Asn 135		: Ile Gly Ile I	
TTA GTA TTT ACT TTA Leu Val Phe Thr Leu 150		ı Asp Ser Glu G	
TTT AGT TAT GAT ATA Phe Ser Tyr Asp Ile 165			
TTT GTA ACT GTT ACT Phe Val Thr Val Thr 180			
AAT GGA AAA TTA ATA Asn Gly Lys Leu Ile 200			

AAT Asn	TTT	AGC Ser	Lys 215	Thr	ATA Ile	ACA Thr	TTT	GAA Glu 220	Ile	AAT Asn	AAA Lys	ATT	CCA Pro 225	Asp	ACC Thr	788
GGT Gly	TTG Leu	ATT Ile 230	Thr	TCA Ser	GAT Asp	TCT	GAT Asp 235	Asn	ATC Ile	AAT Asn	ATG Met	TGG Trp 240	ATA Ile	AGA Arg	GAT Asp	836
TTT Phe	TAT Tyr 245	Ile	TTT Phe	GCT Ala	AAA Lys	GAA Glu 250	TTA Leu	GAT Asp	GGT Gly	AAA Lys	GAT Asp 255	ATT	AAT Asn	ATA Ile	TTA Leu	884
TTT Phe 260	AAT Asn	AGC Ser	TTG Leu	CAA Gln	TAT Tyr 265	ACT Thr	AAT Asn	GTT Val	GTA Val	AAA Lys 270	GAT Asp	TAT Tyr	TGG Trp	GGA Gly	AAT Asn 275	932
GAT Asp	TTA Leu	AGA Arg	TAT	AAT Asn 280	Lys	GAA Glu	TAT Tyr	TAT Tyr	ATG Met 285	GTT Val	AAT Asn	ATA Ile	GAT Asp	TAT Tyr 290	TTA Leu	980
AAT Asn	AGA Arg	TAT	ATG Met 295	TAT Tyr	GCG Ala	AAC Asn	TCA Ser	CGA Arg 300	CAA Gln	ATT Ile	GTT Val	TTT Phe	AAT Asn 305	ACA Thr	CGT Arg	1028
AGA Arg	AAT Asn	AAT Asn 310	AAT Asn	GAC Asp	TTC Phe	AAT Asn	GAA Glu 315	GGA Gly	TAT Tyr	AAA Lys	ATT Ile	ATA Ile 320	ATA Ile	AAA Lys	AGA Arg	1076
ATC Ile	AGA Arg 325	GGA Gly	AAT Asn	ACA Thr	AAT Asn	GAT Asp 330	ACT Thr	AGA Arg	GTA Val	CGA Arg	GGA Gly 335	GGA Gly	GAT Asp	ATT Ile	TTA Leu	1124
TAT Tyr 340	TTT Phe	GAT Asp	ATG Met	ACA Thr	ATT Ile 345	AAT Asn	AAC Asn	AAA Lys	GCA Ala	TAT Tyr 350	AAT Asn	TTG Leu	TTT Phe	ATG Met	AAG Lys 355	1172
AAT Asn	GAA Glu	ACT Thr	ATG Met	TAT Tyr 360	GCA Ala	GAT Asp	AAT Asn	CAT His	AGT Ser 365	ACT Thr	GAA Glu	GAT Asp	ATA Ile	TAT Tyr 370	GCT Ala	1220
ATA Ile	GGT Gly	TTA Leu	AGA Arg 375	GAA Glu	CAA Gln	ACA Thr	AAG Lys	GAT Asp 380	ATA Ile	AAT Asn	GAT Asp	AAT Asn	ATT Ile 385	ATA Ile	TTT Phe	1268
CAA Gln	ATA Ile	CAA Gln 390	CCA Pro	ATG Met	AAT Asn	AAT Asn	ACT Thr 395	TAT Tyr	TAT Tyr	TAC Tyr	GCA Ala	TCT Ser 400	CAA Gln	ATA Ile	TTT Phe	1316
AAA Lys	TCA Ser 405	AAT Asn	TTT Phe	AAT Asn	GGA Gly	GAA Glu 410	AAT Asn	ATT Ile	TCT Ser	GGA Gly	ATA Ile 415	TGT Cys	TCA Ser	ATA Ile	GGT Gly	1364
ACT Thr 420	TAT Tyr	CGT Arg	TTT Phe	AGA Arg	CTT Leu 425	GGA Gly	GGT Gly	GAT Asp	TGG Trp	TAT Tyr 430	AGA Arg	CAC His	AAT Asn	TAT Tyr	TTG Leu 435	1412
GTG Val	CCT Pro	ACT Thr	GTG Val	AAG Lys 440	CAA Gln	GGA Gly	AAT Asn	Tyr	GCT Ala 445	TCA Ser	TTA Leu	TTA Leu	GAA Glu	TCA Ser 450	ACA Thr	1460

TCA ACT CAT TGG GGT TTT GTA CCT GTA AGT GAA TAAAAGCTT Ser Thr His Trp Gly Phe Val Pro Val Ser Glu 455 460

(2) INFORMATION FOR SEQ ID NO:62:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 462 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:62:

Met Gly His His His His His His His His Ser Ser Gly His

Ile Glu Gly Arg His Met Ala Ser Met Ala Leu Leu Lys Asp Ile Ile
20 25 30

Asn Glu Tyr Phe Asn Asn Ile Asn Asp Ser Lys Ile Leu Ser Leu Gln
35 40

Asn Arg Lys Asn Thr Leu Val Asp Thr Ser Gly Tyr Asn Ala Glu Val

Ser Glu Glu Gly Asp Val Gln Leu Asn Pro Ile Phe Pro Phe Asp Phe 65 70 75 80

Lys Leu Gly Ser Ser Gly Glu Asp Arg Gly Lys Val Ile Val Thr Gln
85 90 95

Asn Glu Asn Ile Val Tyr Asn Ser Met Tyr Glu Ser Phe Ser Ile Ser 100 105 110

Phe Trp Ile Arg Ile Asn Lys Trp Val Ser Asn Leu Pro Gly Tyr Thr 115 120 125

Ile Ile Asp Ser Val Lys Asn Asn Ser Gly Trp Ser Ile Gly Ile Ile 130 135 140

Ser Asn Phe Leu Val Phe Thr Leu Lys Gln Asn Glu Asp Ser Glu Gln 145 150 155 160

Ser Ile Asn Phe Ser Tyr Asp Ile Ser Asn Asn Ala Pro Gly Tyr Asn 165 170 175

Lys Trp Phe Phe Val Thr Val Thr Asn Asn Met Met Gly Asn Met Lys 180 185 190

Ile Tyr Ile Asn Gly Lys Leu Ile Asp Thr Ile Lys Val Lys Glu Leu 195 200 205

Thr Gly Ile Asn Phe Ser Lys Thr Ile Thr Phe Glu Ile Asn Lys Ile 210 215 220

Pro Asp Thr Gly Leu Ile Thr Ser Asp Ser Asp Asn Ile Asn Met Trp 225 230 235 240

Ile Arg Asp Phe Tyr Ile Phe Ala Lys Glu Leu Asp Gly Lys Asp Ile 245 250 255 Asn Ile Leu Phe Asn Ser Leu Gln Tyr Thr Asn Val Val Lys Asp Tyr

Trp Gly Asn Asp Leu Arg Tyr Asn Lys Glu Tyr Tyr Met Val Asn Ile

Asp Tyr Leu Asn Arg Tyr Met Tyr Ala Asn Ser Arg Gln Ile Val Phe 295

Asn Thr Arg Arg Asn Asn Asn Asp Phe Asn Glu Gly Tyr Lys Ile Ile 315

Ile Lys Arg Ile Arg Gly Asn Thr Asn Asp Thr Arg Val Arg Gly Gly

Asp Ile Leu Tyr Phe Asp Met Thr Ile Asn Asn Lys Ala Tyr Asn Leu

Phe Met Lys Asn Glu Thr Met Tyr Ala Asp Asn His Ser Thr Glu Asp 360 365

Ile Tyr Ala Ile Gly Leu Arg Glu Gln Thr Lys Asp Ile Asn Asp Asn

Ile Ile Phe Gln Ile Gln Pro Met Asn Asn Thr Tyr Tyr Tyr Ala Ser 390

Gln Ile Phe Lys Ser Asn Phe Asn Gly Glu Asn Ile Ser Gly Ile Cys 405

Ser Ile Gly Thr Tyr Arg Phe Arg Leu Gly Gly Asp Trp Tyr Arg His 425

Asn Tyr Leu Val Pro Thr Val Lys Gln Gly Asn Tyr Ala Ser Leu Leu

Glu Ser Thr Ser Thr His Trp Gly Phe Val Pro Val Ser Glu 455

- (2) INFORMATION FOR SEQ ID NO:63:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 32 base pairs

 - (B) TYPE: nucleic acid(C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "DNA"
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:63:

CGCCATGGCT TTATTAAAAG ATATAATTAA TG

	(i	(A) L B) T C) S	ENGT YPE: TRAN	H: 3 nuc DEDN	CTER 2 ba leic ESS: lin	se p aci sin	airs d								
	(ii							ucle c =								
	(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:64	:					
GCA	AGCT	TTT .	ATTC	ACTT.	AC A	GGTA	CAAA	A CC								32
(2)	INF	ORMA	TION	FOR	SEQ	ID	NO : 6	5 :								
	(i	()	A) L B) T C) S	ENGT YPE : TRAN	H: 3 nuc DEDN		base aci dou	pai d	rs							
	(ii) MO	LECU	LE T	YPE:	DNA	(ge	nomi	c)							
	(ix		A) N.	AME/		CDS	3828									
	(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ :	ID N	0:65	:					
								AAT Asn								48
								CCA Pro 25								96
								CAA Gln								144
								AGT Ser								192
ACT Thr 65	TCA Ser	AAG Lys	TAT Tyr	CAA Gln	AGT Ser 70	TAT Tyr	TAT Tyr	GAT Asp	CCT Pro	AGT Ser 75	TAT Tyr	TTA Leu	TCT Ser	ACT Thr	GAT Asp 80	240
								GGG Gly								288
								AAA Lys 105								336

(2) INFORMATION FOR SEQ ID NO:64:

GGT Gly	TCA Ser	CCT Pro 115	Phe	ATG Met	GGA Gly	GAT Asp	TCA Ser 120	Ser	ACG Thr	CCT Pro	GAA Glu	GAT Asp 125	ACA Thr	TTT Phe	GAT Asp	384
TTT Phe	ACA Thr 130	Arg	CAT His	ACT Thr	ACT Thr	AAT Asn 135	Ile	GCA Ala	GTT Val	GAA Glu	AAG Lys 140	Phe	GAA Glu	AAT Asn	GGT Gly	432
AGT Ser 145	Trp	AAA Lys	GTA Val	ACA Thr	AAT Asn 150	Ile	ATA Ile	ACA Thr	CCA Pro	AGT Ser 155	Val	TTG Leu	ATA Ile	TTT Phe	GGA Gly 160	480
CCA Pro	CTT Leu	CCT Pro	AAT Asn	ATA Ile 165	TTA Leu	GAC Asp	TAT Tyr	ACA Thr	GCA Ala 170	TCC Ser	CTT Leu	ACA Thr	TTG Leu	CAA Gln 175	GGA Gly	528
CAA Gln	CAA Gln	TCA Ser	AAT Asn 180	CCA Pro	TCA Ser	TTT Phe	GAA Glu	GGG Gly 185	TTT Phe	GGA Gly	ACA Thr	TTA Leu	TCT Ser 190	ATA Ile	CTA Leu	576
AAA Lys	GTA Val	GCA Ala 195	CCT Pro	GAA Glu	TTT Phe	TTG Leu	TTA Leu 200	ACA Thr	TTT Phe	AGT Ser	GAT Asp	GTA Val 205	ACA Thr	TCT Ser	AAT Asn	624
CAA Gln	AGT Ser 210	TCA Ser	GCT Ala	GTA Val	TTA Leu	GGC Gly 215	AAA Lys	TCT Ser	ATA Ile	TTT Phe	TGT Cys 220	ATG Met	GAT Asp	CCA Pro	GTA Val	672
ATA Ile 225	GCT Ala	TTA Leu	ATG Met	CAT His	GAG Glu 230	TTA Leu	ACA Thr	CAT His	TCT Ser	TTG Leu 235	CAT His	CAA Gln	TTA Leu	TAT Tyr	GGA Gly 240	720
ATA Ile	AAT Asn	ATA Ile	CCA Pro	TCT Ser 245	GAT Asp	AAA Lys	AGG Arg	ATT Ile	CGT Arg 250	CCA Pro	CAA Glņ	GTT Val	AGC Ser	GAG Glu 255	GGA Gly	768
TTT Phe	TTC Phe	TCT Ser	CAA Gln 260	GAT Asp	GGA Gly	CCC Pro	AAC Asn	GTA Val 265	CAA Gln	TTT Phe	GAG Glu	GAA Glu	TTA Leu 270	TAT Tyr	ACA Thr	816
TTT Phe	GGA Gly	GGA Gly 275	TTA Leu	GAT Asp	GTT Val	Glu	ATA Ile 280	ATA Ile	CCT Pro	CAA Gln	ATT Ile	GAA Glu 285	AGA Arg	TCA Ser	CAA Gln	864
TTA Leu	AGA Arg 290	GAA Glu	AAA Lys	GCA Ala	TTA Leu	GGT Gly 295	CAC His	TAT Tyr	AAA Lys	GAT Asp	ATA Ile 300	GCG Ala	AAA Lys	AGA Arg	CTT Leu	912
AAT Asn 305	AAT Asn	ATT Ile	AAT Asn	AAA Lys	ACT Thr 310	ATT Ile	CCT Pro	TCT Ser	Ser	TGG Trp 315	ATT Ile	AGT Ser	AAT Asn	ATA Ile	GAT Asp 320	960
AAA Lys	TAT Tyr	AAA Lys	AAA Lys	ATA Ile 325	TTT Phe	TCT Ser	GAA Glu	AAG Lys	TAT Tyr 330	Asn	TTT Phe	GAT Asp	AAA Lys	GAT Asp 335	AAT Asn	1008
ACA Thr	GGA Gly	Asn	TTT Phe 340	GTT Val	GTA Val	AAT Asn	Ile	GAT Asp 345	AAA Lys	TTC Phe	AAT Asn	AGC Ser	TTA Leu 350	TAT Tyr	TCA Ser	1056

GAC Asp	TTG Leu	ACT Thr 355	Asn	GTT Val	ATG Met	TCA Ser	GAA Glu 360	Val	GTT Val	TAT Tyr	TCT Ser	TCG Ser 365	Gln	TAT	AAT Asn	1104
GTT Val	AAA Lys 370	Asn	AGG Arg	ACT Thr	CAT	TAT Tyr 375	TTT Phe	TCA Ser	AGG Arg	CAT His	TAT Tyr 380	Leu	CCT Pro	GTA Val	TTT Phe	1152
GCA Ala 385	Asn	ATA Ile	TTA Leu	GAT Asp	GAT Asp 390	AAT Asn	ATT Ile	TAT	ACT Thr	ATA Ile 395	AGA Arg	GAT Asp	GGT Gly	TTT Phe	AAT Asn 400	1200
TTA Leu	ACA Thr	AAT Asn	AAA Lys	GGT Gly 405	TTT Phe	AAT Asn	ATA Ile	GAA Glu	AAT Asn 410	TCG Ser	GGT Gly	CAG Gln	AAT	ATA Ile 415	GAA Glu	1248
AGG Arg	AAT Asn	Pro	GCA Ala 420	Leu	CAA Gln	AAG Lys	CTT Leu	AGT Ser 425	TCA Ser	GAA Glu	AGT Ser	GTA Val	GTA Val 430	GAT Asp	TTA Leu	1296
TTT Phe	ACA Thr	AAA Lys 435	GTA Val	TGT Cys	TTA Leu	AGA Arg	TTA Leu 440	ACA Thr	AAA Lys	AAT Asn	AGT Ser	AGA Arg 445	GAT Asp	GAT Asp	TCA Ser	1344
ACA Thr	TGT Cys 450	ATT Ile	AAA Lys	GTT Val	AAA Lys	AAT Asn 455	AAT Asn	AGA Arg	TTA Leu	CCT Pro	TAT Tyr 460	GTA Val	GCT Ala	GAT Asp	AAA Lys	1392
GAT Asp 465	AGC Ser	ATT Ile	TCA Ser	CAA Gln	GAA Glu 470	ATA Ile	TTT Phe	GAA Glu	AAT Asn	AAA Lys 475	ATT Ile	ATT Ile	ACA Thr	GAT Asp	GAG Glu 480	1440
ACT	AAT Asn	GTA Val	CAA Gln	AAT Asn 485	TAT Tyr	TCA Ser	GAT Asp	AAT Asn	TTT Phe 490	TCA Ser	TTA Leu	GAT Asp	GAA Glu	TCT Ser 495	ATT Ile	1488
TTA Leu	GAT Asp	GGG Gly	CAA Gln 500	GTT Val	CCT Pro	ATT Ile	AAT Asn	CCT Pro 505	GAA Glu	ATA Ile	GTA Val	GAT Asp	CCA Pro 510	CTA Leu	TTA Leu	1536
CCC Pro	AAT Asn	GTT Val 515	AAT Asn	ATG Met	GAA Glu	CCT Pro	TTA Leu 520	AAT Asn	CTT Leu	CCA Pro	GGT Gly	GAA Glu 525	GAA Glu	ATA Ile	GTA Val	1584
Phe	Tyr 530	Asp	Asp	Ile	Thr	AAA Lys 535	Tyr	Val	Asp	Tyr	Leu 540	Asn	Ser	Tyr	Tyr	1632
Tyr 545	Leu	Glu	Ser	Gln	Lys 550	TTA Leu	Ser	Asn	Asn	Val 555	Glu	Asn	Ile	Thr	Leu 560	1680
Thr	Thr	Ser	Val	Glu 565	Glu	GCA Ala	Leu	Gly	Tyr 570	Ser	Asn	Lys	Ile	Tyr 575	Thr	1728
TTT Phe	TTA Leu	CCT Pro	AGC Ser 580	TTA Leu	GCT Ala	GAA Glu	Lys	GTG Val 585	AAT Asn	AAA Lys	GGT Gly	Val	CAA Gln 590	GCA Ala	GGT Gly	1776

TTA Leu	TTC Phe	TTA Leu 595	Asn	TGG Trp	GCG Ala	AAT Asn	GAA Glu 600	Val	GTT Val	GAG Glu	GAT Asp	TTT Phe 605	Thr	ACA Thr	AAT Asn	1824
ATT Ile	ATG Met 610	Lys	AAA Lys	GAT Asp	ACA Thr	TTG Leu 615	GAT Asp	AAA Lys	ATA Ile	TCA Ser	GAT Asp 620	Val	TCA Ser	GTA Val	ATA Ile	1872
ATT Ile 625	Pro	TAT	ATA Ile	GGA Gly	CCT Pro 630	Ala	TTA Leu	AAT Asn	ATA Ile	GGA Gly 635	AAT Asn	TCA Ser	GCA Ala	TTA Leu	AGG Arg 640	1920
GGA Gly	AAT Asn	TTT Phe	AAG Lys	CAA Gln 645	GCA Ala	TTT Phe	GCA Ala	ACA Thr	GCT Ala 650	GGT Gly	GTA Val	GCT Ala	TTT Phe	TTA Leu 655	TTA Leu	1968
GAG Glu	GGA Gly	TTT Phe	CCA Pro 660	GAG Glu	TTT Phe	ACT Thr	ATA Ile	CCT Pro 665	GCA Ala	CTC Leu	GGT Gly	GTA Val	TTT Phe 670	ACC Thr	TTT Phe	2016
TAT	AGT Ser	TCT Ser 675	ATT Ile	CAA Gln	GAA Glu	AGA Arg	GAG Glu 680	AAA Lys	ATT Ile	ATT	AAA Lys	ACT Thr 685	ATA Ile	GAA Glu	AAT Asn	2064
TGT Cys	TTG Leu 690	GAA Glu	CAA Gln	AGA Arg	GTT Val	AAG Lys 695	AGA Arg	TGG Trp	AAA Lys	GAT Asp	TCA Ser 700	TAT Tyr	CAA Gln	TGG Trp	ATG Met	2112
GTA Val 705	TCA Ser	AAT Asn	TGG Trp	TTG Leu	TCA Ser 710	AGA Arg	ATT Ile	ACT Thr	ACT Thr	CAA Gln 715	TTT Phe	AAT Asn	CAT His	ATA Ile	AAT Asn 720	2160
TAT Tyr	CAA Gln	ATG Met	TAT Tyr	GAT Asp 725	TCT Ser	TTA Leu	AGT Ser	TAT Tyr	CAG Gln 730	GCA Ala	GAT Asp	GCA Ala	ATC Ile	AAA Lys 735	GCT Ala	2208
AAA Lys	ATA Ile	GAT Asp	TTA Leu 740	GAA Glu	TAT Tyr	AAA Lys	AAA Lys	TAC Tyr 745	TCA Ser	GGA Gly	AGT Ser	GAT Asp	AAA Lys 750	GAA Glu	AAT Asn	2256
ATA Ile	AAA Lys	AGT Ser 755	CAA Gln	GTT Val	GAA Glu	AAT Asn	TTA Leu 760	AAA Lys	AAT Asn	AGT Ser	TTA Leu	GAT Asp 765	GTA Val	AAA Lys	ATT Ile	2304
TCG Ser	GAA Glu 770	Ala	ATG Met	AAT Asn	AAT Asn	ATA Ile 775	AAT Asn	AAA Lys	TTT Phe	ATA Ile	CGA Arg 780	Glu	TGT Cys	TCT Ser	GTA Val	2352
ACA Thr 785	TAC Tyr	TTA Leu	TTT Phe	AAA Lys	AAT Asn 790	ATG Met	CTC Leu	CCT Pro	AAA Lys	GTA Val 795	ATT Ile	GAC Asp	GAA Glu	TTA Leu	AAT Asn 800	2400
AAG Lys	TTT Phe	GAT Asp	TTA Leu	AGA Arg 805	ACT Thr	AAA Lys	ACA Thr	GAA Glu	TTA Leu 810	ATT Ile	AAT Asn	CTT Leu	ATA Ile	GAT Asp 815	AGT Ser	2448
CAT His	AAT Asn	ATT Ile	ATT Ile 820	CTA Leu	GTT Val	GGT Gly	Glu	GTA Val 825	GAT Asp	AGA Arg	TTA Leu	Lys	GCA Ala 830	AÄA Lys	GTA Val	2496

AAT Asn	GAG Glu	AGT Ser 835	TTT	GAA Glu	AAT Asn	ACA Thr	ATG Met 840	CCT Pro	TTT Phe	AAT Asn	ATT Ile	TTT Phe 845	TCA Ser	TAT	ACT		2544
AAT Asn	AAT Asn 850	TCT Ser	TTA Leu	TTA Leu	AAA Lys	GAT Asp 855	ATA	ATT Ile	AAT Asn	GAA Glu	TAT Tyr 860	Phe	AAT Asn	AGT Ser	ATT Ile		2592
AAT Asn 865	GAT Asp	TCA Ser	AAA Lys	ATT Ile	TTG Leu 870	AGC Ser	TTA Leu	CAA Gln	AAC Asn	AAA Lys 875	AAA Lys	AAT Asn	GCT Ala	TTA Leu	GTG Val 880		2640
GAT Asp	ACA Thr	TCA Ser	GGA Gly	TAT Tyr 885	AAT Asn	GCA Ala	GAA Glu	GTG Val	AGG Arg 890	GTA Val	GGA Gly	GAT Asp	AAT Asn	GTT Val 895	CAA Gln		2688
CTT Leu	AAT Asn	ACG Thr	ATA Ile 900	TAT Tyr	ACA Thr	AAT Asn	GAC Asp	TTT Phe 905	AAA Lys	TTA Leu	AGT Ser	AGT Ser	TCA Ser 910	GGA Gly	GAT Asp		2736
AAA Lys	ATT Ile	ATA Ile 915	GTA Val	AAT Asn	TTA Leu	AAT Asn	AAT Asn 920	AAT Asn	ATT Ile	TTA Leu	TAT Tyr	AGC Ser 925	GCT Ala	ATT Ile	TAT Tyr		2784
GAG Glu	AAC Asn 930	TCT Ser	AGT Ser	GTT Val	AGT Ser	TTT Phe 935	TGG Trp	ATT Ile	AAG Lys	ATA Ile	TCT Ser 940	AAA Lys	GAT Asp	TTA Leu	ACT Thr		2832
AAT Asn 945	TCT Ser	CAT His	AAT Asn	GAA Glu	TAT Tyr 950	ACA Thr	ATA Ile	ATT Ile	AAC Asn	AGT Ser 955	ATA Ile	GAA Glu	CAA Gln	AAT Asn	TCT Ser 960		2880
GGG Gly	TGG Trp	AAA Lys	TTA Leu	TGT Cys 965	ATT Ile	AGG Arg	AAT Asn	GGC Gly	AAT Asn 970	ATA Ile	GAA Glu	TGG Trp	ATT Ile	TTA Leu 975	CAA Gln		2928
GAT Asp	GTT Val	AAT Asn	AGA Arg 980	AAG Lys	TAT Tyr	AAA Lys	AGT Ser	TTA Leu 985	ATT Ile	TTT Phe	GAT Asp	TAT Tyr	AGT Ser 990	GAA Glu	TCA Ser		2976
TTA Leu	AGT Ser	CAT His 995	ACA Thr	GGA Gly	TAT Tyr	ACA Thr	AAT Asn 1000	Lys	TGG Trp	TTT Phe	TTT Phe	GTT Val 1005	Thr	ATA Ile	ACT Thr		3024
AAT Asn	AAT Asn 1010	Ile	ATG Met	GGG Gly	TAT Tyr	ATG Met 1015	Lys	CTT Leu	TAT Tyr	ATA Ile	AAT Asn 1020	Gly	GAA Glu	TTA Leu	AAG Lys		3072
CAG Gln 1025	AGT Ser	CAA Gln	AAA Lys	ATT Ile	GAA Glu 1030	Asp	TTA Leu	GAT Asp	GAG Glu	GTT Val 1035	Lys	TTA Leu	GAT Asp	AAA Lys	ACC Thr 1040	٠	3120
ATA Ile	GTA Val	TTT Phe	GGA Gly	ATA Ile 1045	Asp	GAG Glu	AAT Asn	ATA Ile	GAT Asp 1050	Glu	AAT Asn	CAG Gln	ATG Met	CTT Leu 1055	Trp		3168
ATT	AGA Arg	GAT Asp	TTT Phe 1060	Asn	ATT Ile	TTT Phe	TCT Ser	AAA Lys 1065	Glu	TTA Leu	AGT Ser	Asn	GAA Glu 1070	Asp	ATT Ile		3216

ASI	1 116	107	. 1yr	GIU	GIY	' Gin	11e 108	Leu 0	Arg	Asr	ı Val	l Ile 108	Lys 5	Asp	TAT	326	54
TG(Tr	G GGA O Gly 109	ASI	CCT Pro	TTG Leu	AAG Lys	TTT Phe 109	Asp	ACA Thr	GAA Glu	TAT	TAT Tyr 110	: Ile	ATT Ile	AAT Asn	GAT Asp	. 331	L 2
AAT Asn 110	ı ıyr	ATA Ile	GAT Asp	AGG Arg	TAT Tyr 111	Ile	GCA Ala	CCT Pro	GAA Glu	AGT Ser 111	Asn	GTA Val	CTT Leu	GTA Val	CTT Leu 1120	336	0
GTT Val	Arg	TAT	CCA Pro	GAT Asp 112	Arg	TCT Ser	AAA Lys	TTA Leu	TAT Tyr 113	Thr	GGA Gly	AAT Asn	CCT	ATT Ile 113	Thr	340	8
ATT	AAA Lys	TCA Ser	GTA Val 114	Ser	GAT Asp	AAG Lys	AAT Asn	CCT Pro 114	Tyr	AGT Ser	AGA Arg	ATT	TTA Leu 115	Asn	GGA Gly	345	6
GAT Asp	AAT Asn	ATA Ile 115	ATT Ile 5	CTT Leu	CAT His	ATG Met	TTA Leu 1160	Tyr	AAT Asn	AGT Ser	AGG Arg	AAA Lys 116	Tyr	ATG Met	ATA Ile	350	4
ATA Ile	AGA Arg 117	Asp	ACT Thr	GAT Asp	ACA Thr	ATA Ile 1175	Tyr	GCA Ala	ACA Thr	CAA Gln	GGA Gly 118	Gly	GAG Glu	TGT Cys	TCA Ser	355	2
CAA Gln 118	Asn	TGT Cys	GTA Val	TAT Tyr	GCA Ala 1190	Leu	AAA Lys	TTA Leu	CAG Gln	AGT Ser 119	Asn	TTA Leu	GGT Gly	AAT Asn	TAT Tyr 1200	360	0
GGT Gly	ATA Ile	GGT Gly	ATA Ile	TTT Phe 1205	Ser	ATA Ile	AAA Lys	AAT Asn	ATT Ile 1210	Val	TCT Ser	AAA Lys	AAT Asn	AAA Lys 1215	Tyr	3648	8
Cys	ser	GIN	ATT Ile 1220	Phe	Ser	Ser	Phe	Arg 1225	Glu	Asn	Thr	Met	Leu 1230	Leu	Ala	3696	5
GAT Asp	ATA Ile	TAT Tyr 1235	AAA Lys	CCT Pro	TGG Trp	Arg	TTT Phe 1240	Ser	TTT Phe	AAA Lys	AAT Asn	GCA Ala 1245	Tyr	ACG Thr	CCA Pro	3744	1
GTT Val	GCA Ala 1250	val	ACT Thr	AAT Asn	Tyr	GAA Glu 1255	ACA . Thr	AAA Lys	CTA Leu	TTA Leu	TCA Ser 1260	Thr	TCA Ser	TCT Ser	TTT Phe	3792	}
TGG Trp 1265	Lys	TTT Phe	ATT Ile	Ser .	AGG Arg 1270	GAT (CCA (Pro	GGA Gly	Trp	GTA Val 1275	Glu	TAA				3831	•

(2) INFORMATION FOR SEQ ID NO:66:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1276 amino acids
 (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:66:

Met Thr Trp Pro Val Lys Asp Phe Asn Tyr Ser Asp Pro Val Asn Asp Asn Asp Ile Leu Tyr Leu Arg Ile Pro Gln Asn Lys Leu Ile Thr Thr Pro Val Lys Ala Phe Met Ile Thr Gln Asn Ile Trp Val Ile Pro Glu Arg Phe Ser Ser Asp Thr Asn Pro Ser Leu Ser Lys Pro Pro Arg Pro Thr Ser Lys Tyr Gln Ser Tyr Tyr Asp Pro Ser Tyr Leu Ser Thr Asp Glu Gln Lys Asp Thr Phe Leu Lys Gly Ile Ile Lys Leu Phe Lys Arg Ile Asn Glu Arg Asp Ile Gly Lys Lys Leu Ile Asn Tyr Leu Val Val Gly Ser Pro Phe Met Gly Asp Ser Ser Thr Pro Glu Asp Thr Phe Asp Phe Thr Arg His Thr Thr Asn Ile Ala Val Glu Lys Phe Glu Asn Gly 135 Ser Trp Lys Val Thr Asn Ile Ile Thr Pro Ser Val Leu Ile Phe Gly Pro Leu Pro Asn Ile Leu Asp Tyr Thr Ala Ser Leu Thr Leu Gln Gly 170 Gln Gln Ser Asn Pro Ser Phe Glu Gly Phe Gly Thr Leu Ser Ile Leu 180 185 Lys Val Ala Pro Glu Phe Leu Leu Thr Phe Ser Asp Val Thr Ser Asn 200 Gln Ser Ser Ala Val Leu Gly Lys Ser Ile Phe Cys Met Asp Pro Val Ile Ala Leu Met His Glu Leu Thr His Ser Leu His Gln Leu Tyr Gly 230 235 Ile Asn Ile Pro Ser Asp Lys Arg Ile Arg Pro Gln Val Ser Glu Gly Phe Phe Ser Gln Asp Gly Pro Asn Val Gln Phe Glu Glu Leu Tyr Thr Phe Gly Gly Leu Asp Val Glu Ile Ile Pro Gln Ile Glu Arg Ser Gln 280 Leu Arg Glu Lys Ala Leu Gly His Tyr Lys Asp Ile Ala Lys Arg Leu Asn Asn Ile Asn Lys Thr Ile Pro Ser Ser Trp Ile Ser Asn Ile Asp 305 310 315

Lys Tyr Lys Lys Ile Phe Ser Glu Lys Tyr Asn Phe Asp Lys Asp Asn Thr Gly Asn Phe Val Val Asn Ile Asp Lys Phe Asn Ser Leu Tyr Ser 345 Asp Leu Thr Asn Val Met Ser Glu Val Val Tyr Ser Ser Gln Tyr Asn Val Lys Asn Arg Thr His Tyr Phe Ser Arg His Tyr Leu Pro Val Phe 375 Ala Asn Ile Leu Asp Asp Asn Ile Tyr Thr Ile Arg Asp Gly Phe Asn Leu Thr Asn Lys Gly Phe Asn Ile Glu Asn Ser Gly Gln Asn Ile Glu 410 Arg Asn Pro Ala Leu Gln Lys Leu Ser Ser Glu Ser Val Val Asp Leu Phe Thr Lys Val Cys Leu Arg Leu Thr Lys Asn Ser Arg Asp Asp Ser 440 Thr Cys Ile Lys Val Lys Asn Asn Arg Leu Pro Tyr Val Ala Asp Lys Asp Ser Ile Ser Gln Glu Ile Phe Glu Asn Lys Ile Ile Thr Asp Glu 470 475 Thr Asn Val Gln Asn Tyr Ser Asp Asn Phe Ser Leu Asp Glu Ser Ile Leu Asp Gly Gln Val Pro Ile Asn Pro Glu Ile Val Asp Pro Leu Leu 500 505 Pro Asn Val Asn Met Glu Pro Leu Asn Leu Pro Gly Glu Glu Ile Val Phe Tyr Asp Asp Ile Thr Lys Tyr Val Asp Tyr Leu Asn Ser Tyr Tyr Tyr Leu Glu Ser Gln Lys Leu Ser Asn Asn Val Glu Asn Ile Thr Leu Thr Thr Ser Val Glu Glu Ala Leu Gly Tyr Ser Asn Lys Ile Tyr Thr 565 Phe Leu Pro Ser Leu Ala Glu Lys Val Asn Lys Gly Val Gln Ala Gly Leu Phe Leu Asn Trp Ala Asn Glu Val Val Glu Asp Phe Thr Thr Asn Ile Met Lys Lys Asp Thr Leu Asp Lys Ile Ser Asp Val Ser Val Ile 615 Ile Pro Tyr Ile Gly Pro Ala Leu Asn Ile Gly Asn Ser Ala Leu Arg

Gly Asn Phe Lys Gln Ala Phe Ala Thr Ala Gly Val Ala Phe Leu Leu

645

Glu Gly Phe Pro Glu Phe Thr Ile Pro Ala Leu Gly Val Phe Thr Phe 660 665 670

Tyr Ser Ser Ile Gln Glu Arg Glu Lys Ile Ile Lys Thr Ile Glu Asn 675 680 685

Cys Leu Glu Gln Arg Val Lys Arg Trp Lys Asp Ser Tyr Gln Trp Met 690 695 700

Val Ser Asn Trp Leu Ser Arg Ile Thr Thr Gln Phe Asn His Ile Asn 705 710 715 720

Tyr Gln Met Tyr Asp Ser Leu Ser Tyr Gln Ala Asp Ala Ile Lys Ala 725 730 735

Lys Ile Asp Leu Glu Tyr Lys Lys Tyr Ser Gly Ser Asp Lys Glu Asn
740 745 750

Ile Lys Ser Gln Val Glu Asn Leu Lys Asn Ser Leu Asp Val Lys Ile
755 760 765

Ser Glu Ala Met Asn Asn Ile Asn Lys Phe Ile Arg Glu Cys Ser Val 770 775 780

Thr Tyr Leu Phe Lys Asn Met Leu Pro Lys Val Ile Asp Glu Leu Asn 785 790 795 800

Lys Phe Asp Leu Arg Thr Lys Thr Glu Leu Ile Asn Leu Ile Asp Ser 805 810 815

His Asn Ile Ile Leu Val Gly Glu Val Asp Arg Leu Lys Ala Lys Val 820 825 830

Asn Glu Ser Phe Glu Asn Thr Met Pro Phe Asn Ile Phe Ser Tyr Thr 835 840 845

Asn Asn Ser Leu Leu Lys Asp Ile Ile Asn Glu Tyr Phe Asn Ser Ile 850 855 860

Asn Asp Ser Lys Ile Leu Ser Leu Gln Asn Lys Lys Asn Ala Leu Val 865 870 875 880

Asp Thr Ser Gly Tyr Asn Ala Glu Val Arg Val Gly Asp Asn Val Gln 885 890 895

Leu Asn Thr Ile Tyr Thr Asn Asp Phe Lys Leu Ser Ser Ser Gly Asp 900 905 910

Lys Ile Ile Val Asn Leu Asn Asn Ile Leu Tyr Ser Ala Ile Tyr 915 920 925

Glu Asn Ser Ser Val Ser Phe Trp Ile Lys Ile Ser Lys Asp Leu Thr 930 935 940

Asn Ser His Asn Glu Tyr Thr Ile Ile Asn Ser Ile Glu Gln Asn Ser 945 950 955 960

Gly Trp Lys Leu Cys Ile Arg Asn Gly Asn Ile Glu Trp Ile Leu Gln 965 970 975

Asp Val Asn Arg Lys Tyr Lys Ser Leu Ile Phe Asp Tyr Ser Glu Ser 980 985 990 Leu Ser His Thr Gly Tyr Thr Asn Lys Trp Phe Phe Val Thr Ile Thr 995 1000 1005

- Asn Asn Ile Met Gly Tyr Met Lys Leu Tyr Ile Asn Gly Glu Leu Lys 1010 1015 1020
- Gln Ser Gln Lys Ile Glu Asp Leu Asp Glu Val Lys Leu Asp Lys Thr 1025 1030 1035 1040
- Ile Val Phe Gly Ile Asp Glu Asn Ile Asp Glu Asn Gln Met Leu Trp
 1045 1050 1055
- Ile Arg Asp Phe Asn Ile Phe Ser Lys Glu Leu Ser Asn Glu Asp Ile 1060 1065 1070
- Asn Ile Val Tyr Glu Gly Gln Ile Leu Arg Asn Val Ile Lys Asp Tyr 1075 1080 1085
- Trp Gly Asn Pro Leu Lys Phe Asp Thr Glu Tyr Tyr Ile Ile Asn Asp 1090 1095 1100
- Asn Tyr Ile Asp Arg Tyr Ile Ala Pro Glu Ser Asn Val Leu Val Leu 1105 1110 1115 1120
- Val Arg Tyr Pro Asp Arg Ser Lys Leu Tyr Thr Gly Asn Pro Ile Thr 1125 1130 1135
- Ile Lys Ser Val Ser Asp Lys Asn Pro Tyr Ser Arg Ile Leu Asn Gly
 1140 1145 1150
- Asp Asn Ile Ile Leu His Met Leu Tyr Asn Ser Arg Lys Tyr Met Ile 1155 1160 1165
- Ile Arg Asp Thr Asp Thr Ile Tyr Ala Thr Gln Gly Gly Glu Cys Ser
- Gln Asn Cys Val Tyr Ala Leu Lys Leu Gln Ser Asn Leu Gly Asn Tyr 1185 1190 1195 1200
- Gly Ile Gly Ile Phe Ser Ile Lys Asn Ile Val Ser Lys Asn Lys Tyr 1205 1210 1215
- Cys Ser Gln Ile Phe Ser Ser Phe Arg Glu Asn Thr Met Leu Leu Ala 1220 1225 1230
- Asp Ile Tyr Lys Pro Trp Arg Phe Ser Phe Lys Asn Ala Tyr Thr Pro 1235 1240 1245
- Val Ala Val Thr Asn Tyr Glu Thr Lys Leu Leu Ser Thr Ser Ser Phe 1250 1255 1260
- Trp Lys Phe Ile Ser Arg Asp Pro Gly Trp Val Glu 1265 1270 1275

(2)	INF	ORMA	TION	FOR	SEC	ID	NO : 6	7:								
	(i	(A) L B) T C) S	ENGT YPE : TRAN	H: 1 nuc DEDN	CTER 469 leic ESS: lin	base aci dou	pai d	.rs							
	(ii) MO	LECU	LE T	YPE:	DNA	(ge	nomi	c)							
	(ix		A) N	AME/		CDS 108		60								
	(xi) SE	QUEN	CE D	ESCR	IPTI	ON:	SEQ	ID N	0:67	:			٠		
AGA	TCTC	GAT	CCCG	CGAA	AT T	AATA	CGAC	T CA	CTAT	AGGG	GAA	TTGT	GAG	CGGA	TAACAA	60
TTC	CCCT	CTA	gaaa'	TAAT	TT T	GTTT.	AACT	т та	AGAA	GGAG	ATA	TACC		GGC		116
CAT His	CAT His 5	CAT His	CAT His	CAT His	CAT His	CAT His 10	CAT His	CAC His	AGC Ser	AGC Ser	GGC Gly 15	CAT His	ATC Ile	GAA Glu	GGT Gly	164
CGT Arg 20	CAT His	ATG Met	GCT Ala	AGC Ser	ATG Met 25	GCT Ala	TTA Leu	TTA Leu	AAA Lys	GAT Asp 30	ATA Ile	ATT Ile	AAT Asn	GAA Glu	TAT Tyr 35	212
TTC Phe	AAT Asn	AGT Ser	ATT Ile	AAT Asn 40	GAT Asp	TCA Ser	AAA Lys	ATT Ile	TTG Leu 45	AGC Ser	TTA Leu	CAA Gln	AAC Asn	AAA Lys 50	AAA Lys	260
AAT Asn	GCT Ala	TTA Leu	GTG Val 55	GAT Asp	ACA Thr	TCA Ser	GGA Gly	TAT Tyr 60	AAT Asn	GCA Ala	GAA Glu	GTG Val	AGG Arg 65	GTA Val	GGA Gly	308
GAT Asp	AAT Asn	GTT Val 70	CAA Gln	CTT Leu	AAT Asn	ACG Thr	ATA Ile 75	TAT Tyr	ACA Thr	AAT Asn	GAC Asp	TTT Phe 80	AAA Lys	TTA .Leu	AGT Ser	356
AGT Ser	TCA Ser 85	Gly	Asp	Lys	Ile	ATA Ile 90	Val	AAT Asn	TTA Leu	AAT Asn	AAT Asn 95	AAT Asn	ATT Ile	TTA Leu	TAT Tyr	404
AGC Ser 100	GCT Ala	ATT Ile	TAT Tyr	GAG Glu	AAC Asn 105	TCT Ser	AGT Ser	GTT Val	AGT Ser	TTT Phe 110	TGG Trp	ATT Ile	AAG Lys	ATA Ile	TCT Ser 115	452
AAA Lys	GAT Asp	TTA Leu	ACT Thr	AAT Asn 120	TCT Ser	CAT His	AAT Asn	GAA Glu	TAT Tyr 125	ACA Thr	ATA Ile	ATT Ile	AAC Asn	AGT Ser 130	ATA Ile	500
										AGG Arg						548
										AAA Lys						596

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ТА Ту	T AG r Se 16	- 61	A TC u Se	A TT	A AG' u Se:	r CAT	5 Th	A GG	A ТА У Ту:	r AC.	A AA r As 17	n Ly:	A TG	G TT p Ph	T TTT e Phe	644
GT Va 18		r AT.	A AC' e Th	r Aa:	r AA: n Asi 185	1 TTE	A ATO	G GGG G Gly	TA'	r ATG	t Ly	A CT? s Lev	TA:	r Ata	A AAT e Asn 195	692
GGI Gl	A GAJ y Glu	TT	A AAG 1 Lys	G CAC Glr 200	ı ser	CAA Glm	AAA Lys	ATT Ile	GA/ Glu 205	ı Ası	r TT/	A GAT 1 Asp	GAC Glu	GT: 1 Val 210	r AAG l Lys D	740
TT) Let	A GAT 1 Asp	Lys	A ACC 5 Thr 215	: ile	GTA Val	TTT.	GGA Gly	ATA Ile 220	: Asp	GAC Glu	AA1 Asr	T ATA	GAT Asp 225	Gli	G AAT 1 Asn	788
Glr	ATC Met	Leu 230	rrth	ATT Ile	AGA Arg	GAT Asp	TTT Phe 235	Asn	ATT	TTI Phe	TCT Ser	AAA Lys 240	Glu	Leu	A AGT	836
AAT Asn	GAA Glu 245	wah	T ATT	AAT Asn	ATT	GTA Val 250	TAT	GAG Glu	GGA Gly	CAA Gln	ATA Ile 255	Leu	AGA Arg	AAT	GTT Val	884
ATT Ile 260	пÀ2	GAT Asp	TAT Tyr	TGG Trp	GGA Gly 265	AAT Asn	CCT Pro	TTG Leu	AAG Lys	TTT Phe 270	GAT Asp	ACA Thr	GAA Glu	TAT Tyr	TAT Tyr 275	932
ATT	ATT	AAT Asn	GAT Asp	AAT Asn 280	TAT	ATA Ile	GAT Asp	AGG Arg	TAT Tyr 285	ATT Ile	GCA Ala	CCT Pro	GAA Glu	AGT Ser 290	AAT Asn	980
GTA Val	CTT Leu	GTA Val	CTT Leu 295	GTT Val	CGG Arg	TAT Tyr	CCA Pró	GAT Asp 300	AGA Arg	TCT	AAA Lys	TTA Leu	TAT Tyr 305	ACT Thr	GGA Gly	1028
AAT Asn	CCT Pro	ATT Ile 310	ACT Thr	ATT Ile	AAA Lys	TCA Ser	GTA Val 315	TCT Ser	GAT Asp	AAG Lys	AAT Asn	CCT Pro 320	TAT Tyr	AGT Ser	AGA Arg	1076
ATT Ile	TTA Leu 325	AAT Asn	GGA Gly	GAT Asp	AAT Asn	ATA Ile 330	ATT Ile	CTT Leu	CAT His	ATG Met	TTA Leu 335	TAT Tyr	AAT Asn	AGT Ser	AGG Arg	1124
AAA Lys 340	TAT Tyr	ATG Met	ATA Ile	ATA Ile	AGA Arg 345	GAT Asp	ACT Thr	GAT Asp	ACA Thr	ATA Ile 350	TAT Tyr	GCA Ala	ACA Thr	CAA Gln	GGA Gly 355	1172
GGA Gly	GAG Glu	TGT Cys	TCA Ser	CAA Gln 360	AAT Asn	TGT Cys	GTA Val	Tyr	GCA Ala 365	TTA Leu	AAA Lys	TTA Leu	CAG Gln	AGT Ser 370	AAT Asn	1220
TTA Leu	GGT Gly	AAT Asn	TAT Tyr 375	GGT Gly	ATA Ile	GGT . Gly	Ile	TTT . Phe 380	AGT Ser	ATA Ile	AAA Lys	Asn	ATT Ile 385	GTA Val	TCT Ser	1268
AAA Lys	ASI	AAA Lys 390	TAT Tyr	TGT Cys	AGT Ser	CAA /	ATT Ile 395	TTC 'Phe	TCT . Ser	AGT Ser	Phe	AGG (Arg (GAA Glu	AAT Asn	ACA Thr	1316

ATG CTT CTA GCA GAT ATA TAT AAA CCT TGG AGA TTT TCT TTT AAA AAT Met Leu Leu Ala Asp Ile Tyr Lys Pro Trp Arg Phe Ser Phe Lys Asn 405
GCA TAC ACG CCA GTT GCA GTA ACT AAT TAT GAA ACA AAA CTA TTA TCA Ala Tyr Thr Pro Val Ala Val Thr Asn Tyr Glu Thr Lys Leu Leu Ser 420 430 435
ACT TCA TCT TTT TGG AAA TTT ATT TCT AGG GAT CCA GGA TGG GTA GAG Thr Ser Ser Phe Trp Lys Phe Ile Ser Arg Asp Pro Gly Trp Val Glu 440 445 450
TAAAAGCTT
(2) INFORMATION FOR SEQ ID NO:68:
(i) SEQUENCE CHARACTERISTICS: (A) LENGTH: 451 amino acids (B) TYPE: amino acid (D) TOPOLOGY: linear
(ii) MOLECULE TYPE: protein
(xi) SEQUENCE DESCRIPTION: SEQ ID NO:68:
Met Gly His His His His His His His His His Ser Ser Gly His 1 10 15
Ile Glu Gly Arg His Met Ala Ser Met Ala Leu Leu Lys Asp Ile Ile 20 25 30
Asn Glu Tyr Phe Asn Ser Ile Asn Asp Ser Lys Ile Leu Ser Leu Gln 35 40 45
Asn Lys Lys Asn Ala Leu Val Asp Thr Ser Gly Tyr Asn Ala Glu Val 50 60
Arg Val Gly Asp Asn Val Gln Leu Asn Thr Ile Tyr Thr Asn Asp Phe 65 70 75 80
Lys Leu Ser Ser Ser Gly Asp Lys Ile Ile Val Asn Leu Asn Asn Asn 90 95
Ile Leu Tyr Ser Ala Ile Tyr Glu Asn Ser Ser Val Ser Phe Trp Ile 100 105 110
Lys Ile Ser Lys Asp Leu Thr Asn Ser His Asn Glu Tyr Thr Ile Ile 115 120 125
Asn Ser Ile Glu Gln Asn Ser Gly Trp Lys Leu Cys Ile Arg Asn Gly 130 135 140
Asn Ile Glu Trp Ile Leu Gln Asp Val Asn Arg Lys Tyr Lys Ser Leu 145 150 155 160

Ile Phe Asp Tyr Ser Glu Ser Leu Ser His Thr Gly Tyr Thr Asn Lys

Trp Phe Phe Val Thr Ile Thr Asn Asn Ile Met Gly Tyr Met Lys Leu

Tyr Ile Asn Gly Glu Leu Lys Gln Ser Gln Lys Ile Glu Asp Leu Asp 195 200 205

Glu Val Lys Leu Asp Lys Thr Ile Val Phe Gly Ile Asp Glu Asn Ile 210 215 220

Asp Glu Asn Gln Met Leu Trp Ile Arg Asp Phe Asn Ile Phe Ser Lys 225 230 235 240

Glu Leu Ser Asn Glu Asp Ile Asn Ile Val Tyr Glu Gly Gln Ile Leu 245 250 255

Arg Asn Val Ile Lys Asp Tyr Trp Gly Asn Pro Leu Lys Phe Asp Thr 260 265 270

Glu Tyr Tyr Ile Ile Asn Asp Asn Tyr Ile Asp Arg Tyr Ile Ala Pro 275 280 285

Glu Ser Asn Val Leu Val Leu Val Arg Tyr Pro Asp Arg Ser Lys Leu 290 295 300

Tyr Thr Gly Asn Pro Ile Thr Ile Lys Ser Val Ser Asp Lys Asn Pro 305 310 315 320

Tyr Ser Arg Ile Leu Asn Gly Asp Asn Ile Ile Leu His Met Leu Tyr 325 330 335

Asn Ser Arg Lys Tyr Met Ile Ile Arg Asp Thr Asp Thr Ile Tyr Ala 340 345 350

Thr Gln Gly Glu Cys Ser Gln Asn Cys Val Tyr Ala Leu Lys Leu 355 360 365

Gln Ser Asn Leu Gly Asn Tyr Gly Ile Gly Ile Phe Ser Ile Lys Asn 370 375 380

Ile Val Ser Lys Asn Lys Tyr Cys Ser Gln Ile Phe Ser Ser Phe Arg 385 390 395 400

Glu Asn Thr Met Leu Leu Ala Asp Ile Tyr Lys Pro Trp Arg Phe Ser 405 410 415

Phe Lys Asn Ala Tyr Thr Pro Val Ala Val Thr Asn Tyr Glu Thr Lys
420 425 430

Leu Leu Ser Thr Ser Ser Phe Trp Lys Phe Ile Ser Arg Asp Pro Gly
435 440 445

Trp Val Glu 450

(2) INFORMATION FOR SEQ ID NO:69:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 32 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: other nucleic acid
 (A) DESCRIPTION: /desc = "DNA"

GCA	AGCT	TTT	ACTC	TACC	CA T	CCTG	GATC	C CT								32
(2)	INF	orma	TION	FOR	SEQ	ID	NO : 7	0:								
	(i	(QUEN A) L B) T C) S D) T	ENGT YPE: TRAN	H: 3 nuc DEDN	825 leic ESS:	base aci dou	pai d	rs							
	(ii) MO	LECU	LE T	YPE:	DNA	(ge	nomi	c)							
	(ix	(.	ATUR A) N B) L	AME/			3822									
	(xi) SE	QUEN	CE D	ESCR	IPTI(ON:	SEQ	ID N	0:70	:					
													GTT Val			48
													AGT Ser 30		AAA Lys	96
													ATT Ile			144
													CCG Pro		TCA Ser	192
													TTA Leu			240
													TTA Leu			288
	Ile	Asn		Asn	Pro	Ala	Gly	Lys	Val	Leu	Leu	Gln	GAA Glu 110	Ile	TCA Ser	336
													GAT Asp			384
													TCA Ser			432
													GCA Ala			480

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:69:

GAT Asp	ATA Ile	TTT Phe	GAA Glu	AGT Ser 165	TGT Cys	TGT Cys	TAC Tyr	CCC Pro	GTT Val 170	AGA Arg	AAA Lys	CTA Leu	ATA Ile	GAT Asp 175	CCA Pro	528
GAT Asp	GTA Val	GTT Val	TAT Tyr 180	GAT Asp	CCA Pro	AGT Ser	AAT Asn	TAT Tyr 185	GGT Gly	TTT Phe	GGA Gly	TCA Ser	ATT Ile 190	AAT Asn	ATC Ile	576
GTG Val	ACA Thr	TTT Phe 195	TCA Ser	CCT Pro	GAG Glu	TAT Tyr	GAA Glu 200	TAT Tyr	ACT Thr	TTT Phe	AAT Asn	GAT Asp 205	ATT Ile	AGT Ser	GGA Gly	624
					ACA Thr											672
					ATA Ile 230											720
					GAG Glu											768
					ATA Ile											816
					ATT Ile											864
					TAT Tyr											912
					GAA Glu 310											960
					CTA Leu											1008
					AAT Asn											1056
					AAT Asn											1104
					TTT Phe											1152
					GAG Glu 390											1200

AAT Asn	CGC	GGA Gly	CAA Glm	AG' Sei 405		A AAG e Lys	G TT S Le	A AA u As	T CC n Pr 41	о гу	A AT	T AT	T GA e As	T TO P Se 41	C ATT r Ile 5	1248
CCA Pro	GAT Asp	AAA Lys	GGT Gly 420		A GTA	A GAZ L Glu	A AA	G AT s Il 42	e va	T AA. 1 Ly:	A TT s Ph	T TG1 e Cys	T AAG 5 Lys 430	s Se	C GTT r Val	1296
ATT Ile	CCT Pro	AGA Arg 435	~, 5	GGT Gly	ACA Thi	A AAG Lys	GCC Ala	a Pro	A CC	G CG	A CT	A TGC u Cys 445	Ile	r AG	A GTA g Val	1344
	AAT Asn 450	AGT Ser	GAG Glu	TTA Leu	TT1 Phe	Phe 455	val	A GC:	r TC	A GAZ r Glu	A AG: 1 Se: 460	Ser	TAT Tyr	AA' Ası	r GAA n Glu	1392
AAT Asn 465	GAT Asp	ATT Ile	AAT Asn	ACA Thr	CCT Pro 470	пås	GAA Glu	ATT	GAC Asp	GAT Asp 475	Thr	ACA Thr	AAT Asn	CT/ Let	A AAT ASN 480	1440
AAT . Asn .	AAT Asn	TAT Tyr	AGA Arg	AAT Asn 485	AAT Asn	TTA Leu	GAT Asp	GAA Glu	GTI Val 490	Ile	TTA Leu	GAT Asp	TAT Tyr	AAT Asn 495		1488
CAG I	ACA Thr	ATA Ile	CCT Pro 500	CAA Gln	ATA Ile	TCA Ser	AAT Asn	CGA Arg 505	Thr	. TTA Leu	AAT Asn	ACA Thr	CTT Leu 510	GTA Val	CAA Gln	1536
GAC A		AGT Ser 515	TAT Tyr	GTG Val	CCA Pro	AGA Arg	TAT Tyr 520	GAT Asp	TCT Ser	AAT Asn	GGA Gly	ACA Thr 525	AGT Ser	GAA Glu	ATA Ile	1584
GAG G	SAA : Slu :	TAT Tyr	GAT Asp	GTT Val	GTT Val	GAC Asp 535	TTT Phe	AAT Asn	GTA Val	TTT	TTC Phe 540	TAT Tyr	TTA Leu	CAT His	GCA Ala	1632
CAA A Gln L 545	AA (ys l	GTG (CCA (Pro (3 L U	GGT Gly 550	GAA Glu	ACC Thr	AAT Asn	ATA Ile	AGT Ser 555	TTA Leu	ACT Thr	TCT Ser	TCA Ser	ATT Ile 560	1680
GAT A Asp T	CA G	SCA 1	ne a 1	TTA Leu 565	GAA Glu	GAA Glu	TCC Ser	AAA Lys	GAT Asp 570	ATA Ile	TTT Phe	TTT Phe	TCT Ser	TCA Ser 575	GAG Glu	1728
TTT A	TC G le A	י קבי	ACT F Thr 1 580	TC I	AAT . Asn	AAA (Lys)	Pro	GTA Val 585	AAT Asn	GCA Ala	GCA Ala	Leu	TTT Phe 590	ATA Ile	GAT Asp	1776
TGG A'		GC A er L 95	AA G	TA /	ATA . Ile .	arg A	SAT Asp 500	TTT Phe	ACC Thr	ACT Thr	Glu	GCT 1 Ala 1 605	ACA (CAA Gln	AAA Lys	1824
AGT AG Ser Th	CT G ar V. 10	TT G al A	AT A sp L	AG A	rie 1	GCA (Ala <i>A</i> 515	SAC A	ATA Ile	TCT Ser	Leu	ATT Ile 620	GTA (Val I	CCC :	FAT Fyr	GTA Val	1872
GGT CT Gly Le 625	PT Ge eu Al	CT T la L	TG A eu A	211 1	TA A le 1	ATT A [le I	TT (GAG (Ala (GAA 1 Glu 1 635	AAA (Lys (GGA A	AT I	he	GAG Glu 640	1920

GA(G GCA 1 Ala	A TTT	GAA Glu	Leu 645	Let	GGA Gly	GT(G GG:	T ATT	Leu	TT/	A GAZ 1 Glu	A TT:	GT(Va) 65	G CCA l Pro 5	1968
GAA Glu	A CTI	C ACA	ATT Ile 660	Pro	GTA Val	ATT	TTA Lev	GT(Va) 665	Phe	ACG Thr	ATA Ile	A AAA E Lys	TC0 Ser 670	Ty	r ATA	2016
GAT Asp	TCA Ser	TAT Tyr 675	GIU	AAT Asn	AAA Lys	AAT Asn	AAA Lys 680	Ala	ATT	AAA Lys	GCA Ala	ATA Ile 685	Asr	AA1 Asr	TCA Ser	2064
TTA Leu	ATC Ile 690	Glu	AGA Arg	GAA Glu	GCA Ala	AAG Lys 695	TGG Trp	AAA Lys	GAA Glu	ATA Ile	TAT Tyr 700	Ser	TGG	ATA Ile	GTA Val	2112
TCA Ser 705	Asn	TGG Trp	CTT Leu	ACT Thr	AGA Arg 710	Ile	AAT Asn	ACT Thr	CAA Gln	TTT Phe 715	AAT Asn	AAA Lys	AGA Arg	AAA Lys	GAG Glu 720	2160
CAA Gln	ATG Met	TAT Tyr	CAG Gln	GCT Ala 725	TTA Leu	CAA Gln	AAT Asn	CAA Gln	GTA Val 730	GAT Asp	GCA Ala	ATA Ile	AAA Lys	ACA Thr 735	GCA Ala	2208
ATA Ile	GAA Glu	TAT Tyr	AAA Lys 740	TAT Tyr	AAT Asn	AAT Asn	TAT Tyr	ACT Thr 745	TCA Ser	GAT Asp	GAG Glu	AAA Lys	AAT Asn 750	AGA Arg	CTT	2256
GAA Glu	TCT	GAA Glu 755	TAT	AAT Asn	ATC Ile	AAT Asn	AAT Asn 760	ATA Ile	GAA Glu	GAA Glu	GAA Glu	TTG Leu 765	AAT Asn	AAA Lys	AAA Lys	2304
GTT Val	TCT Ser 770	TTA Leu	GCA Ala	ATG Met	AAA Lys	AAT Asn 775	Ile	GAA Glu	AGA Arg	TTT Phe	ATG Met 780	ACA Thr	GAA Glu	AGT Ser	TCT Ser	2352
ATA Ile 785	TCT Ser	TAT Tyr	TTA Leu	ATG Met	AAA Lys 790	TTA Leu	ATA Ile	AAT Asn	GAA Glu	GCC Ala 795	AAA Lys	GTT Val	GGT Gly	AAA Lys	TTA Leu 800	2400
AAA Lys	AAA Lys	TAT Tyr	GAT Asp	AAC Asn 805	His	GTT Val	AAG Lys	AGC Ser	GAT Asp 810	TTA Leu	TTA Leu	AAC Asn	TAT Tyr	ATT Ile 815	CTC Leu	2448
GAC Asp	CAT His	AGA Arg	TCA Ser 820	ATC Ile	TTA Leu	GGA Gly	GAG Glu	CAG Gln 825	ACA Thr	AAT Asn	GAA Glu	TTA Leu	AGT Ser 830	GAT Asp	TTG Leu	2496
GTG Val	ACT Thr	AGT Ser 835	ACT Thr	TTG Leu	AAT Asn	Ser	AGT Ser 840	ATT Ile	CCA Pro	TTT Phe	GAA Glu	CTT Leu 845	TCT Ser	TCA Ser	TAT Tyr	2544
ACT Thr	AAT Asn 850	GAT Asp	AAA Lys	ATT Ile	Leu	ATT Ile 855	ATA Ile	TAT Tyr	TTT Phe	Asn	AGA Arg 860	TTA Leu	TAT Tyr	AAA Lys	AAA Lys	2592
ATT Ile 865	AAA Lys	GAT Asp	AGT Ser	Ser	ATT Ile 870	TTA (Leu)	GAT Asp	ATG Met	Arg	TAT Tyr 875	GAA Glu	AAT . Asn .	AAT Asn	AAA Lys	TTT Phe 880	2640

AT.	A GA e As	T AT	C TC	r GG	A TA	r GGT	TC	A AA	r ATA	A AGO	C AT	T AA 1	GGA	AAC	GTA	2688
				883	•	r Gly			890)				895	;	
TA' Ty:	r Il	T TA:	r TCA Ser 900	Ini	A AA: C Asi	AGA Arg	AAT Asn	CAA Glr 905	n Phe	GG/	A ATA	TAT Tyr	AAT Asn 910	Ser	' AGG ' Arg	2736
CT. Let	r AG' 1 Sei	r GAZ r Glu 915	ıvaı	AAT Asn	TATA	GCT Ala	CAA Gln 920	Asn	TAAT Asn	GAT Asp	T ATT	ATA Ile 925	TAC	AAT Asn	AGT Ser	2784
AG! Arg	TAT TY1 930	GII	AAT Asn	TTI Phe	AG1 Ser	ATT Ile 935	AGT Ser	TTC Phe	TGG	GTA Val	AGG Arg 940	Ile	CCT Pro	AAA Lys	CAC His	2832
TAC Tyr 945	- uys	CCT Pro	ATG Met	AAT Asn	CAT His 950	AAT Asn	CGG Arg	GAA Glu	TAC	ACT Thr 955	Ile	ATA	AAT Asn	TGT Cys	ATG Met 960	2880
GGG Gly	AAT Asn	AAT Asn	AAT Asn	TCG Ser 965	GGA Gly	TGG Trp	AAA Lys	ATA Ile	TCA Ser 970	CTT Leu	AGA Arg	ACT Thr	GTT Val	AGA Arg 975	GAT Asp	2928
TGT Cys	GAA Glu	ATA Ile	ATT Ile 980	TGG Trp	ACT Thr	TTA Leu	CAA Gln	GAT Asp 985	ACT Thr	TCT Ser	GGA Gly	AAT Asn	AAG Lys 990	GAA Glu	AAT Asn	2976
TTA Leu	ATT	TTT Phe 995	AGG Arg	TAT Tyr	GAA Glu	GAA Glu	CTT Leu 1000	Asn	AGG Arg	ATA Ile	TCT	AAT Asn 1005	Tyr	ATA Ile	AAT Asn	3024
AAA Lys	TGG Trp 101	TIE	TTT Phe	GTA Val	ACT Thr	ATT Ile 1015	Thr	AAT Asn	AAT Asn	AGA Arg	TTA Leu 1020	Gly	AAT Asn	TCT Ser	AGA Arg	3072
ATT Ile 1029	IYI	ATC Ile	AAT Asn	GGA Gly	AAT Asn 1030	TTA Leu	ATA Ile	GTT Val	GAA Glu	AAÁ Lys 1035	Ser	ATT Ile	TCG Ser	Asn	TTA Leu 1040	3120
GGT Gly	GAT Asp	ATT	CAT His	GTT Val 1045	Ser	GAT . Asp .	AAT Asn	ATA Ile	TTA Leu 1050	Phe	AAA Lys	ATT (Val	GGT Gly 1055	TGT Cys	3168
GAT Asp	GAT Asp	GAA Glu	ACG Thr 1060	- 1 -	GTT Val	GGT A	TTC.	AGA Arg 1065	TAT	TTT Phe	AAA Lys	vaT 1	TTT / Phe /	AAT / Asn '	ACG Thr	3216
GAA Glu	TTA Leu	GAT Asp 1075	Lys	ACA Thr	GAA Glu	ATT (GAG Glu 1080	ACT Thr	TTA '	TAT . Tyr	Ser .	AAT (Asn (GAG (Glu 1	CCA (Pro 1	GAT Asp	3264
CCA Pro	AGT Ser 1090	TTE	TTA . Leu :	AAA Lys	Asn	TAT 7 Tyr 7	rgg (rp (GGA . Gly .	AAT ' Asn '	Tyr :	TTG (Leu)	CTA 1	TAT /	AAT / Asn 1	AAA Lys	3312
AAA Lys 1105	TAT	TAT Tyr	TTA '	rue '	AAT ' Asn '	TTA (Leu I	CTA I	AGA A	Lys i	GAT A Asp 1	AAG 1 Lys 1	TAT A Tyr I	ATT A	Thr I	CTG Leu 1120	3360

	ATT TTA AAT A Ile Leu Asn I 1125				3408
Ser Val Phe	TTG AAC TAT A Leu Asn Tyr I 1140	Glu Gly V		Ile Ile	3456
•	GGT CCT ATA G				3504
	CTA GCA TAC A Leu Ala Tyr I	Val Asp A			3552
	GCT GAT ACA A Ala Asp Thr I 1190				3600
	AAC GAT AGC T Asn Asp Ser I 1205		/al Met Asp		3648
Gly Asn Asn	TGC ACA ATG A Cys Thr Met A 1220	Asn Asn A		Asn Ile	3696
	GGT TTT CAT T Gly Phe His S	 			3744
	ATA CGA AGA A Ile Arg Arg A 1	Ser Asn G			3792
	AAA GAG AAT G Lys Glu Asn G 1270			:	3825

- (2) INFORMATION FOR SEQ ID NO:71:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1274 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: protein
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:71:

Met Pro Val Ala Ile Asn Ser Phe Asn Tyr Asn Asp Pro Val Asn Asp 1 5 10 15

Asp Thr Ile Leu Tyr Met Gln Ile Pro Tyr Glu Glu Lys Ser Lys Lys 20 25 30

Tyr Tyr Lys Ala Phe Glu Ile Met Arg Asn Val Trp Ile Ile Pro Glu 35 40 45

Arg Asn Thr Ile Gly Thr Asn Pro Ser Asp Phe Asp Pro Pro Ala Ser 50 55 60

Leu Lys Asn Gly Ser Ser Ala Tyr Tyr Asp Pro Asn Tyr Leu Thr Thr Asp Ala Glu Lys Asp Arg Tyr Leu Lys Thr Thr Ile Lys Leu Phe Lys 90 Arg Ile Asn Ser Asn Pro Ala Gly Lys Val Leu Leu Gln Glu Ile Ser Tyr Ala Lys Pro Tyr Leu Gly Asn Asp His Thr Pro Ile Asp Glu Phe Ser Pro Val Thr Arg Thr Thr Ser Val Asn Ile Lys Leu Ser Thr Asn 135 Val Glu Ser Ser Met Leu Leu Asn Leu Leu Val Leu Gly Ala Gly Pro 150 Asp Ile Phe Glu Ser Cys Cys Tyr Pro Val Arg Lys Leu Ile Asp Pro Asp Val Val Tyr Asp Pro Ser Asn Tyr Gly Phe Gly Ser Ile Asn Ile Val Thr Phe Ser Pro Glu Tyr Glu Tyr Thr Phe Asn Asp Ile Ser Gly 200 Gly His Asn Ser Ser Thr Glu Ser Phe Ile Ala Asp Pro Ala Ile Ser Leu Ala His Glu Leu Ile His Ala Leu His Gly Leu Tyr Gly Ala Arg Gly Val Thr Tyr Glu Glu Thr Ile Glu Val Lys Gln Ala Pro Leu Met 250 Ile Ala Glu Lys Pro Ile Arg Leu Glu Glu Phe Leu Thr Phe Gly Gly Gln Asp Leu Asn Ile Ile Thr Ser Ala Met Lys Glu Lys Ile Tyr Asn 280 Asn Leu Leu Ala Asn Tyr Glu Lys Ile Ala Thr Arg Leu Ser Glu Val 290 295 Asn Ser Ala Pro Pro Glu Tyr Asp Ile Asn Glu Tyr Lys Asp Tyr Phe Gln Trp Lys Tyr Gly Leu Asp Lys Asn Ala Asp Gly Ser Tyr Thr Val Asn Glu Asn Lys Phe Asn Glu Ile Tyr Lys Lys Leu Tyr Ser Phe Thr 345 Glu Ser Asp Leu Ala Asn Lys Phe Lys Val Lys Cys Arg Asn Thr Tyr Phe Ile Lys Tyr Glu Phe Leu Lys Val Pro Asn Leu Leu Asp Asp Asp Ile Tyr Thr Val Ser Glu Gly Phe Asn Ile Gly Asn Leu Ala Val Asn 395

Asn Arg Gly Gln Ser Ile Lys Leu Asn Pro Lys Ile Ile Asp Ser Ile Pro Asp Lys Gly Leu Val Glu Lys Ile Val Lys Phe Cys Lys Ser Val Ile Pro Arg Lys Gly Thr Lys Ala Pro Pro Arg Leu Cys Ile Arg Val Asn Asn Ser Glu Leu Phe Phe Val Ala Ser Glu Ser Ser Tyr Asn Glu 455 Asn Asp Ile Asn Thr Pro Lys Glu Ile Asp Asp Thr Thr Asn Leu Asn Asn Asn Tyr Arg Asn Asn Leu Asp Glu Val Ile Leu Asp Tyr Asn Ser 490 . Gln Thr Ile Pro Gln Ile Ser Asn Arg Thr Leu Asn Thr Leu Val Gln 500 Asp Asn Ser Tyr Val Pro Arg Tyr Asp Ser Asn Gly Thr Ser Glu Ile Glu Glu Tyr Asp Val Val Asp Phe Asn Val Phe Phe Tyr Leu His Ala Gln Lys Val Pro Glu Gly Glu Thr Asn Ile Ser Leu Thr Ser Ser Ile 555 Asp Thr Ala Leu Leu Glu Glu Ser Lys Asp Ile Phe Phe Ser Ser Glu 565 Phe Ile Asp Thr Ile Asn Lys Pro Val Asn Ala Ala Leu Phe Ile Asp Trp Ile Ser Lys Val Ile Arg Asp Phe Thr Thr Glu Ala Thr Gln Lys Ser Thr Val Asp Lys Ile Ala Asp Ile Ser Leu Ile Val Pro Tyr Val Gly Leu Ala Leu Asn Ile Ile Ile Glu Ala Glu Lys Gly Asn Phe Glu Glu Ala Phe Glu Leu Leu Gly Val Gly Ile Leu Leu Glu Phe Val Pro Glu Leu Thr Ile Pro Val Ile Leu Val Phe Thr Ile Lys Ser Tyr Ile 665 Asp Ser Tyr Glu Asn Lys Asn Lys Ala Ile Lys Ala Ile Asn Asn Ser Leu Ile Glu Arg Glu Ala Lys Trp Lys Glu Ile Tyr Ser Trp Ile Val Ser Asn Trp Leu Thr Arg Ile Asn Thr Gln Phe Asn Lys Arg Lys Glu

Gln Met Tyr Gln Ala Leu Gln Asn Gln Val Asp Ala Ile Lys Thr Ala

Ile Glu Tyr Lys Tyr Asn Asn Tyr Thr Ser Asp Glu Lys Asn Arg Leu
740 745 750

Glu Ser Glu Tyr Asn Ile Asn Asn Ile Glu Glu Glu Leu Asn Lys Lys
755 760 765

Val Ser Leu Ala Met Lys Asn Ile Glu Arg Phe Met Thr Glu Ser Ser 770 775 780

Ile Ser Tyr Leu Met Lys Leu Ile Asn Glu Ala Lys Val Gly Lys Leu 785 790 795 800

Lys Lys Tyr Asp Asn His Val Lys Ser Asp Leu Leu Asn Tyr Ile Leu 805 810 815

Asp His Arg Ser Ile Leu Gly Glu Gln Thr Asn Glu Leu Ser Asp Leu 820 825 830

Val Thr Ser Thr Leu Asn Ser Ser Ile Pro Phe Glu Leu Ser Ser Tyr 835 840 845

Thr Asn Asp Lys Ile Leu Ile Ile Tyr Phe Asn Arg Leu Tyr Lys Lys 850 855 860

Ile Lys Asp Ser Ser Ile Leu Asp Met Arg Tyr Glu Asn Asn Lys Phe 865 870 875 880

Ile Asp Ile Ser Gly Tyr Gly Ser Asn Ile Ser Ile Asn Gly Asn Val 885 890 895

Tyr Ile Tyr Ser Thr Asn Arg Asn Gln Phe Gly Ile Tyr Asn Ser Arg 900 905 910

Leu Ser Glu Val Asn Ile Ala Gln Asn Asn Asp Ile Ile Tyr Asn Ser 915 920 925

Arg Tyr Gln Asn Phe Ser Ile Ser Phe Trp Val Arg Ile Pro Lys His 930 935 940

Tyr Lys Pro Met Asn His Asn Arg Glu Tyr Thr Ile Ile Asn Cys Met 945 950 955 960

Gly Asn Asn Asn Ser Gly Trp Lys Ile Ser Leu Arg Thr Val Arg Asp 965 970 975

Cys Glu Ile Ile Trp Thr Leu Gln Asp Thr Ser Gly Asn Lys Glu Asn 980 985 990

Leu Ile Phe Arg Tyr Glu Glu Leu Asn Arg Ile Ser Asn Tyr Ile Asn 995 1000 1005

Lys Trp Ile Phe Val Thr Ile Thr Asn Asn Arg Leu Gly Asn Ser Arg 1010 1015 1020

Ile Tyr Ile Asn Gly Asn Leu Ile Val Glu Lys Ser Ile Ser Asn Leu 1025 1030 1035 1040

Gly Asp Ile His Val Ser Asp Asn Ile Leu Phe Lys Ile Val Gly Cys 1045 1050 1055

Asp Asp Glu Thr Tyr Val Gly Ile Arg Tyr Phe Lys Val Phe Asn Thr 1060 1065 1070

Pro	Ser 109		Leu	Lys	Asn	Tyr 109		Gly	Asn	Туг	Leu 110		Tyr	Asn	Lys		
Lys 110		Tyr	Leu	Phe	Asn 111		Leu	Arg	Lys	Asp 1115	Lys 5	Tyr	Ile	Thr	Leu 1120		
Asn	Ser	Gly	Ile	Leu 1125		Ile	Asn		Gln 1130	_	Gly	Val	Thr	Glu 1139	_		
Ser	Val	Phe	Leu 1140		Tyr	Lys	Leu	Tyr 1149		Gly	Val	Glu	Val 1150		Ile		
Arg	Lys	Asn 115		Pro	Ile	Asp	Ile 1160		Asn	Thr	Asp	Asn 1165		Val	Arg		
Lys	Asn 1170	_	Leu	Ala	Tyr	Ile 1179		Val	Val	Asp	Arg 1180		Val	Glu	Tyr		
Arg 1185		Tyr	Ala	Asp	Thr 1190	-	Ser	Glu	Lys	Glu 1199	Lys	Ile	Ile	Arg	Thr 1200		
Ser	Asn	Leu	Asn	Asp 1205		Leu	Gly	Gln	Ile 1210		Val	Met	Asp	Ser 1215			
Gly	Asn	Asn	Cys 1220		Met	Asn	Phe	Gln 1225		Asn	Asn	Gly	Ser 1230		Ile		
Gly	Leu	Leu 1235	_	Phe	His	Ser	Asn 1240		Leu	Val	Ala	Ser 1245		Trp	Tyr		
Tyr	Asn 1250		Ile	Arg	Arg	Asn 1255		Ser	Ser	Asn	Gly 1260	_	Phe	Trp	Ser		
Ser 1265		Ser	Lys	Glu	Asn 1270	_	Trp	Lys	Glu								
(2)	INFO	RMA	rion	FOR	SEQ	ID N	10:72	: :									
	(i)	(Z (E	A) LE B) TY C) ST	ength (PE : [ran]	I: 14 nucl	60 b	STIC ase acid doub ar	pair l	s		,						
	(ii)	MOI	LECUI	LE TY	PE:	DNA	(ger	omic	:)								
	(ix)	(2	ATURE A) NA B) LO	ME/F			.145	51									
	(xi)	SEÇ	QUENC	CE DE	SCRI	PTIC	N: S	EQ I	D NC	: 72 :							
AGAT	CTCC	SAT C	CCGC	GAAA	T TA	ATAC	GACT	CAC	TATA	GGG	GAAT	TGTG	AG C	GGAT	AACAA		60
TTCC	CCTC	CTA C	TAAA	TAATI	T TO	TTT	ACTI	TAA	GAAG	GAG	ATAT	ACC	ATG Met			1	16

Glu Leu Asp Lys Thr Glu Ile Glu Thr Leu Tyr Ser Asn Glu Pro Asp 1075 1080 1085

CAT His	CAT His 5	His	CAT His	CAT His	CAT His	CAT His 10	CAT His	CAC	AGC Ser	AGC Ser	GGC Gly 15	His	ATC Ile	GAA Glu	GGT Gly	164
CGT Arg 20	CAT His	ATG Met	GCT Ala	AGC Ser	ATG Met 25	Ala	ATT Ile	CTA Leu	ATT	ATA Ile 30	Tyr	TTT Phe	AAT Asn	AGA Arg	TTA Leu 35	212
TAT Tyr	AAA Lys	AAA Lys	ATT Ile	AAA Lys 40	Asp	AGT Ser	TCT Ser	ATT	TTA Leu 45	GAT Asp	ATG Met	CGA Arg	TAT Tyr	GAA Glu 50	AAT Asn	260
AAT Asn	AAA Lys	TTT	ATA Ile 55	GAT Asp	ATC Ile	TCT	GGA Gly	TAT Tyr 60	GGT Gly	TCA Ser	AAT Asn	ATA Ile	AGC Ser 65	ATT Ile	AAT Asn	308
GGA Gly	AAC Asn	GTA Val 70	TAT Tyr	ATT Ile	TAT Tyr	TCA Ser	ACA Thr 75	AAT Asn	AGA Arg	AAT Asn	CAA Gln	TTT Phe 80	GGA Gly	ATA Ile	TAT Tyr	356
AAT Asn	AGT Ser 85	AGG Arg	CTT Leu	AGT Ser	GAA Glu	GTT Val 90	AAT Asn	ATA Ile	GCT Ala	CAA Gln	AAT Asn 95	AAT Asn	GAT Asp	ATT	ATA Ile	404
TAC Tyr 100	AAT Asn	AGT Ser	AGA Arg	TAT Tyr	CAA Gln 105	AAT Asn	TTT Phe	AGT Ser	ATT Ile	AGT Ser 110	TTC Phe	TGG Trp	GTA Val	AGG Arg	ATT Ile 115	452
CCT Pro	AAA Lys	CAC His	TAC Tyr	AAA Lys 120	CCT Pro	ATG Met	AAT Asn	CAT His	AAT Asn 125	CGG Arg	GAA Glu	TAC Tyr	Thr	ATA Ile 130	ATA Ile	500
AAT Asn	TGT Cys	ATG Met	GGG Gly 135	AAT Asn	AAT Asn	AAT Asn	TCG Ser	GGA Gly 140	TGG Trp	AAA Lys	ATA Ile	TCA Ser	CTT Leu 145	AGA Arg	ACT Thr	548
GTT Val	AGA Arg	GAT Asp 150	TGT Cys	GAA Glu	ATA Ile	ATT Ile	TGG Trp 155	ACT Thr	TTA Leu	CAA Gln	GAT Asp	ACT Thr 160	TCT Ser	GGA Gly	AAT Asn	596
AAG Lys	GAA Glu 165	AAT Asn	TTA Leu	ATT Ile	TTT Phe	AGG Arg 170	TAT Tyr	GAA Glu	GAA Glu	CTT Leu	AAT Asn 175	AGG Arg	ATA Ile	TCT Ser	AAT Asn	644
TAT Tyr 180	ATA Ile	AAT Asn	AAA Lys	TGG Trp	ATT Ile 185	TTT Phe	GTA Val	ACT Thr	ATT Ile	ACT Thr 190	AAT Asn	AAT Asn	AGA Arg	TTA Leu	GGC Gly 195	692
AAT Asn	TCT Ser	AGA Arg	ATT Ile	TAC Tyr 200	ATC Ile	AAT Asn	GGA Gly	AAT Asn	TTA Leu 205	ATA Ile	GTT Val	GAA Glu	AAA Lys	TCA Ser 210	ATT Ile	740
TCG Ser	AAT Asn	TTA Leu	GGT Gly 215	GAT Asp	ATT Ile	CAT His	GTT Val	AGT Ser 220	GAT Asp	AAT Asn	ATA Ile	TTA Leu	TTT Phe 225	AAA Lys	ATT Ile	788
GTT Val	GGT Gly	TGT Cys 230	GAT Asp	GAT Asp	GAA Glu	Thr	TAT Tyr 235	GTT Val	GGT Gly	ATA Ile	AGA Arg	TAT Tyr 240	TTT Phe	AAA Lys	GTT Val	836

TTT Phe	AAT Asn 245	ACG Thr	GAA Glu	TTA Leu	GAT Asp	AAA Lys 250	ACA Thr	GAA Glu	ATT Ile	GAG Glu	ACT Thr 255	TTA Leu	TAT	AGT Ser	AAT Asn	884
						TTA Leu										932
						TTA Leu										980
						ATT Ile										1028
						TTG Leu										1076
						GGT Gly 330										1124
						CTA Leu										1172
						GCT Ala										1220
						AAC Asn										1268
						TGC Cys										1316
						GGT Gly 410						_				1364
						ATA Ile										1412
_						AAA Lys							TGAA	AGCT	T	1460

(2) INFORMATION FOR SEQ ID NO:73:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 448 amino acids
 (B) TYPE: amino acid
 (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:73:

Met Gly His His His His His His His His His Ser Ser Gly His

1 10 15

Ile Glu Gly Arg His Met Ala Ser Met Ala Ile Leu Ile Ile Tyr Phe 20 25 30

Asn Arg Leu Tyr Lys Lys Ile Lys Asp Ser Ser Ile Leu Asp Met Arg
35 40

Tyr Glu Asn Asn Lys Phe Ile Asp Ile Ser Gly Tyr Gly Ser Asn Ile
50 55

Ser Ile Asn Gly Asn Val Tyr Ile Tyr Ser Thr Asn Arg Asn Gln Phe 65 70 75 80

Gly Ile Tyr Asn Ser Arg Leu Ser Glu Val Asn Ile Ala Gln Asn Asn 85 90 95

Asp Ile Ile Tyr Asn Ser Arg Tyr Gln Asn Phe Ser Ile Ser Phe Trp
100 105 110

Val Arg Ile Pro Lys His Tyr Lys Pro Met Asn His Asn Arg Glu Tyr 115 120 125

Thr Ile Ile Asn Cys Met Gly Asn Asn Asn Ser Gly Trp Lys Ile Ser 130 140

Leu Arg Thr Val Arg Asp Cys Glu Ile Ile Trp Thr Leu Gln Asp Thr 145 150 155 160

Ser Gly Asn Lys Glu Asn Leu Ile Phe Arg Tyr Glu Glu Leu Asn Arg 165 170 175

Ile Ser Asn Tyr Ile Asn Lys Trp Ile Phe Val Thr Ile Thr Asn Asn 180 185 190

Arg Leu Gly Asn Ser Arg Ile Tyr Ile Asn Gly Asn Leu Ile Val Glu 195 200 205

Lys Ser Ile Ser Asn Leu Gly Asp Ile His Val Ser Asp Asn Ile Leu 210 215 220

Phe Lys Ile Val Gly Cys Asp Asp Glu Thr Tyr Val Gly Ile Arg Tyr 225 230 235 240

Phe Lys Val Phe Asn Thr Glu Leu Asp Lys Thr Glu Ile Glu Thr Leu 245 250 255

Tyr Ser Asn Glu Pro Asp Pro Ser Ile Leu Lys Asn Tyr Trp Gly Asn 260 265 270

Tyr Leu Leu Tyr Asn Lys Lys Tyr Tyr Leu Phe Asn Leu Leu Arg Lys 275 280 285

Asp Lys Tyr Ile Thr Leu Asn Ser Gly Ile Leu Asn Ile Asn Gln Gln
290 295 300

Arg Gly Val Thr Glu Gly Ser Val Phe Leu Asn Tyr Lys Leu Tyr Glu 305 310 315 320

Gly	Val	Glu	Val	Ile 325	Ile	Arg	Lys	Asn	Gly 330	Pro	Ile	Asp	Ile	Ser 335	Asn		
Thr	Asp	Asn	Phe 340	Val	Arg	Lys	Asn	Asp 345	Leu	Ala	Tyr	Ile	Asn 350	Val	Val		
Asp	Arg	Gly 355	Val	Glu	Tyr	Arg	Leu 360	Tyr	Ala	Asp	Thr	Lys 365	Ser	Glu	Lys		
Glu	Lys 370	Ile	Ile	Arg	Thr	Ser 375	Asn	Leu	Asn	Asp	Ser 380	Leu	Gly	Gln	Ile		
Ile 385	Val	Met	Asp	Ser	Ile 390	Gly	Asn	Asn	Cys	Thr 395	Met	Asn	Phe	Gln	Asn 400		
Asn	Asn	Gly	Ser	Asn 405	Ile	Gly	Leu	Leu	Gly 410	Phe	His	Ser	Asn	Asn 415	Leu		
Val	Ala	Ser	Ser 420	Trp	Tyr	Tyr	Asn	Asn 425	Ile	Arg	Arg	Asn	Thr 430	Ser	Ser		
Asn	Gly	Cys 435	Phe	Trp	Ser	Ser	Ile 440	Ser	Lys	Glu	Asn	Gly 445	Trp	Lys	Glu		
(2)	INFO	RMAT	CION	FOR	SEQ	ID N	iO:74	:				•					
		(B (C (D	L) LE S) TY C) ST O) TO	NGTH PE: RAND POLO	: 33 nucl EDNE GY:	bas eic SS: line	e pa acid sing ar	irs l									
		(A	.) DE	SCRI	PTIO	N: /	desc	clei	DNA"								
	(xi)	SEQ	UENC	E DE	SCRI	PTIO	N: S	EQ I	D NO	:74:							
CGCC	ATGG	CT A	TTCT	AATT	А ТА	TATT	TTAA	TAG									33
(2)	INFO	RMAT	ION	FOR	SEQ	ID N	0:75	:									
	(i)	(B) LE) TY) ST	NGTH PE: RAND	: 29 nucl EDNE	TERI: base eic a SS:	e pa acid sing	irs								·	
	(ii)							clei		iđ							
	(xi)	SEQ	UENC:	E DE	SCRI	PTIO	N: S	EQ I	D NO	:75:							
GCAA	GCTT'	TC A	TTCT	TTCC	A TC	CATT	CTC	:									29

	()		(A) I (B) 1	LENG? TYPE: TRAN	TH: 3 DEDM	894 Cleic NESS:	RISTI base aci dou lear	pa:	irs							
	(ii	.) MC	LECU	TE 1	YPE:	DNA	(ge	imon	.c)							
		(AME/ OCAI	'ION :	1	3891						-			
አ ጥር							ON:									
Met 1	Pro	Val	Asn	Ile 5	Lys	Asn	Phe	Asn Asn	TAT Tyr 10	Asn	.GAC	Pro	I ATT	AAT Asn 15	AAT Asn	4.8
GAT Asp	GAC Asp	ATT	ATT Ile 20	Met	ATG Met	GAA Glu	CCA Pro	TTC Phe 25	AAT Asn	GAC Asp	CCA Pro	GGG Gly	CCA Pro	Gly	ACA Thr	96
TAT	TAT	AAA Lys 35	Ala	TTT Phe	AGG Arg	ATT Ile	ATA Ile 40	GAT Asp	CGT Arg	ATT Ile	TGG Trp	ATA Ile 45	Val	CCA Pro	GAA Glu	144
AGG Arg	TTT Phe 50	Thr	TAT	GGA Gly	TTT Phe	CAA Gln 55	Pro	GAC Asp	CAA Gln	TTT Phe	AAT Asn 60	GCC Ala	AGT Ser	ACA Thr	GGA Gly	192
GTT Val 65	TTT Phe	AGT Ser	AAA Lys	GAT Asp	GTC Val	TAC Tyr	GAA Glu	TAT Tyr	TAC Tyr	GAT Asp 75	CCA Pro	ACT Thr	TAT Tyr	TTA Leu	AAA Lys 80	240
ACC Thr	GAT Asp	GCT Ala	GAA Glu	AAA Lys 85	GAT Asp	AAA Lys	TTT Phe	TTA Leu	AAA Lys 90	ACA Thr	ATG Met	ATT Ile	AAA Lys	TTA Leu 95	TTT Phe	288
AAT Asn	AGA Arg	ATT Ile	AAT Asn 100	TCA Ser	AAA Lys	CCA Pro	TCA Ser	GGA Gly 105	CAG Gln	AGA Arg	TTA Leu	CTG Leu	GAT Asp 110	ATG Met	ATA Ile	336
GTA Val	GAT Asp	GCT Ala 115	ATA Ile	CCT Pro	TAT Tyr	CTT Leu	GGA Gly 120	AAT Asn	GCA Ala	TCT Ser	ACA Thr	CCG Pro 125	CCC Pro	GAC Asp	AAA Lys	384
TTT Phe	GCA Ala 130	GCA Ala	AAT Asn	GTT Val	GCA Ala	AAT Asn 135	GTA Val	TCT Ser	ATT Ile	AAT Asn	AAA Lys 140	AAA Lys	ATT Ile	ATC Ile	CAA Gln	432
CCT Pro 145	GGA Gly	GCT Ala	GAA Glu	GAT Asp	CAA Gln 150	ATA Ile	AAA Lys	GGT Gly	TTA Leu	ATG Met 155	ACA Thr	AAT Asn	TTA Leu	ATA Ile	ATA Ile 160	480
TTT Phe	GGA Gly	CCA Pro	GGA Gly	CCA Pro 165	GTT Val	CTA Leu	AGT Ser	GAT Asp	AAT Asn 170	TTT Phe	ACT Thr	GAT Asp	AGT Ser	ATG Met 175	ATT Ile	528

(2) INFORMATION FOR SEQ ID NO:76:

ATG Met	AAT Asn	GGC Gly	CAT His 180	TCC Ser	CCA Pro	ATA Ile	TCA Ser	GAA Glu 185	GGA Gly	TTT Phe	GGT Gly	GCA Ala	AGA Arg 190	ATG Met	ATG Met		576
	AGA Arg														GAA Glu		624
	AAA Lys 210														CCA Pro		672
_	CTA Leu					Glu	-										720
	ATT Ile																768
	ATG Met																816
	GGA Gly									-							864
	AAT Asn 290																912
	GTT Val																960
	ATA Ile																1008
	AGT Ser																1056
	GGC Gly																1104
	TAT Tyr 370																1152
	TTA Leu																1200
	AAA Lys															٠.	1248

AAA Lys	GAG Glu	GCT Ala	TAT Tyr 420	GAA Glu	GAA Glu	ATC Ile	AGC Ser	CTA Leu 425	GAA Glu	CAT His	CTC Leu	GTT Val	ATA Ile 430	TAT Tyr	AGA Arg	1296
ATA Ile	GCA Ala	ATG Met 435	TGC Cys	AAG Lys	CCT Pro	GTA Val	ATG Met 440	TAC Tyr	AAA Lys	AAT Asn	ACC Thr	GGT Gly 445	AAA Lys	TCT Ser	GAA Glu	1344
CAG Gln	TGT Cys 450	ATT Ile	ATT Ile	GTT Val	AAT Asn	AAT Asn 455	GAG Glu	GAT Asp	TTA Leu	TTT Phe	TTC Phe 460	ATA Ile	GCT Ala	AAT Asn	AAA Lys	1392
GAT Asp 465	AGT Ser	TTT Phe	TCA Ser	AAA Lys	GAT Asp 470	TTA Leu	GCT Ala	AAA Lys	GCA Ala	GAA Glu 475	ACT Thr	ATA Ile	GCA Ala	TAT Tyr	AAT Asn 480	1440
ACA Thr	CAA Gln	AAT Asn	AAT Asn	ACT Thr 485	ATA Ile	GAA Glu	AAT Asn	AAT Asn	TTT Phe 490	TCT Ser	ATA Ile	GAT Asp	CAG Gln	TTG Leu 495	ATT Ile	1488
TTA Leu	GAT Asp	AAT Asn	GAT Asp 500	TTA Leu	AGC Ser	AGT Ser	GGC Gly	ATA Ile 505	GAC Asp	TTA Leu	CCA Pro	AAT Asn	GAA Glu 510	AAC Asn	ACA Thr	1536
GAA Glu	CCA Pro	TTT Phe 515	ACA Thr	AAT Asn	TTT Phe	GAC Asp	GAC Asp 520	ATA Ile	GAT Asp	ATC Ile	CCT Pro	GTG Val 525	TAT Tyr	ATT Ile	AAA Lys	1584
CAA Gln	TCT Ser 530	GCT Ala	TTA Leu	AAA Lys	AAA Lys	ATT Ile 535	TTT Phe	GTG Val	GAT Asp	GGA Gly	GAT Asp 540	AGC Ser	CTT Leu	TTT Phe	GAA Glu	1632
TAT Tyr 545	TTA Leu	CAT	GCT Ala	CAA Gln	ACA Thr 550	TTT Phe	CCT Pro	TCT Ser	AAT Asn	ATA Ile 555	GAA Glu	AAT Asn	CTA Leu	CAA Gln	CTA Leu 560	1680
ACG Thr	AAT Asn	TCA Ser	TTA Leu	AAT Asn 565	GAT Asp	GCT Ala	TTA Leu	AGA Arg	AAT Asn 570	AAT Asn	AAT Asn	AAA Lys	GTC Val	TAT Tyr 575	ACT Thr	1728
TTT Phe	TTT Phe	TCT Ser	ACA Thr 580	AAC Asn	CTT	GTT Val	GAA Glu	AAA Lys 585	GCT Ala	AAT Asn	ACA Thr	GTT Val	GTA Val 590	GGT Gly	GCT Ala	1776
TCA Ser	CTT Leu	TTT Phe 595	GTA Val	AAC Asn	TGG Trp	GTA Val	AAA Lys 600	GGA Gly	GTA Val	ATA Ile	GAT Asp	GAT Asp 605	TTT Phe	ACA Thr	TCT Ser	1824
GAA Glu	TCC Ser 610	ACA Thr	CAA Gln	AAA Lys	AGT Ser	ACT Thr 615	ATA Ile	GAT Asp	AAA Lys	GTT Val	TCA Ser 620	GAT Asp	GTA Val	TCC Ser	ATA Ile	1872
ATT Ile 625	ATT	CCC Pro	TAT Tyr	ATA Ile	GGA Gly 630	CCT Pro	GCT Ala	TTG Leu	AAT Asn	GTA Val 635	GGA Gly	AAT Asn	GAA Glu	ACA Thr	GCT Ala 640	1920
AAA Lys	GAA Glu	AAT Asn	TTT Phe	AAA Lys 645	AAT Asn	GCT Ala	TTT Phe	GAA Glu	ATA Ile 650	GGT Gly	GGA Gly	GCC Ala	GCT Ala	ATC Ile 655	TTA Leu	1968

			ATT Ile						2016
			AAA Lys 680						2064
			CAA Gln						2112
			ACG Thr					ATA Ile 720	2160
			TTA Leu						2208
			AAT Asn						2256
			GAT Asp 760						2304
			ATA Ile						2352
			ATG Met						2400
			AAG Lys				_		2448
			GAT Asp						2496
			AGT Ser 840						2544
			CAA Gln						2592
			AGT Ser						2640
			GCA Ala						2688

ATC Ile	TTT Phe	AAT Asn	GAT Asp 900	ATA Ile	GGA Gly	AAT Asn	GGT Gly	CAA Gln 905	TTT Phe	AAA Lys	TTA Leu	AAT Asn	AAT Asn 910	TCT Ser	GAA Glu	.2736
AAT Asn	AGT Ser	AAT Asn 915	ATT Ile	ACG Thr	GCA Ala	CAT His	CAA Gln 920	AGT Ser	AAA Lys	TTC Phe	GTT Val	GTA Val 925	TAT	GAT Asp	AGT Ser	2784
ATG Met	TTT Phe 930	GAT Asp	AAT Asn	TTT Phe	AGC Ser	ATT Ile 935	AAC Asn	TTT Phe	TGG Trp	GTA Val	AGG Arg 940	ACT Thr	CCT	AAA Lys	TAT Tyr	2832
AAT Asn 945	AAT Asn	AAT Asn	GAT Asp	ATA Ile	CAA Gln 950	ACT Thr	TAT Tyr	CTT Leu	CAA Gln	AAT Asn 955	GAG Glu	TAT Tyr	ACA Thr	ATA Ile	ATT Ile 960	2880
												ATT Ile				2928
												TCT Ser				2976
TTT Phe	TTC Phe	GAA Glu 995	TAT Tyr	AGT Ser	ATA Ile	AAA Lys	GAT Asp 1000	Asn	ATA Ile	TCA Ser	GAT Asp	TAT Tyr 1009	Ile	AAT Asn	AAA Lys	3024
Trp	Phe 1010	Ser	Ile	Thr	Ile	Thr 1015	Asn	Asp	Arg	Leu	Gly 1020		Ala	Asn	Ile	3072
Tyr 1025	Ile	Asn	Gly	Ser	Leu 1030	Lys)	Lys	Ser	Glu	Lys 1035	Ile	TTA Leu	Asn	Leu	Asp 1040	3120
Arg	Ile	Asn	Ser	Ser 1045	Asn	Asp	Ile	Asp	Phe 1050	Lys)	Leu	ATT Ile	Asn	Cys 1055	Thr	3168
Asp	Thr	Thr	Lys 1060	Phe	Val	Trp	Ile	Lys 1065	Asp	Phe	Asn	ATT Ile	Phe 1070	Gly)	Arg	3216
Glu	Leu	Asn 1075	Ala	Thr	Glu	Val	Ser 1080	Ser	Leu	Tyr	Trp	ATT Ile 1085	Gln ;	Ser	Ser	3264
Thr	Asn 1090	Thr	Leu	Lys	Asp	Phe 1095	Trp	Gly	Asn	Pro	Leu 1100		Tyr	Asp	Thr	3312
Gln 1105	Tyr	Tyr	Leu	Phe	Asn 1110	Gln	Gly	Met	Gln	Asn 1115	Ile	TAT Tyr	Ile	Lys	Tyr 1120	3360
					Met					Pro		ACA Thr			Asn	3408

				Asn					Tyr			TTA Leu		Phe		3456
			Ala					Asn				GAT Asp 1165	Asn			3504
		Gly					Leu					ATT Ile				3552
	Tyr					Leu					Glu	ATT Ile				3600
					Ile					Thr		TAT Tyr			Leu	3648
				Tyr					Thr			TGT Cys		Ile		3696
			Asp					Gly				ATT Ile 1245	Gly			3744
		Asp					Trp					AAT Asn				3792
	Ser					Arg					Asn	ATA Ile				3840
					Trp					Val		GAA Glu			Thr	3888
GAA Glu	TAA															3894

(2) INFORMATION FOR SEQ ID NO:77:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1297 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:77:

Met Pro Val Asn Ile Lys Asn Phe Asn Tyr Asn Asp Pro Ile Asn Asn 1 5 10 15

Asp Asp Ile Ile Met Met Glu Pro Phe Asn Asp Pro Gly Pro Gly Thr 20 25 30

Tyr Tyr Lys Ala Phe Arg Ile Ile Asp Arg Ile Trp Ile Val Pro Glu 35 40 45

Arg Phe Thr Tyr Gly Phe Gln Pro Asp Gln Phe Asn Ala Ser Thr Gly 55 Val Phe Ser Lys Asp Val Tyr Glu Tyr Tyr Asp Pro Thr Tyr Leu Lys Thr Asp Ala Glu Lys Asp Lys Phe Leu Lys Thr Met Ile Lys Leu Phe Asn Arg Ile Asn Ser Lys Pro Ser Gly Gln Arg Leu Leu Asp Met Ile Val Asp Ala Ile Pro Tyr Leu Gly Asn Ala Ser Thr Pro Pro Asp Lys Phe Ala Ala Asn Val Ala Asn Val Ser Ile Asn Lys Lys Ile Ile Gln Pro Gly Ala Glu Asp Gln Ile Lys Gly Leu Met Thr Asn Leu Ile Ile Phe Gly Pro Gly Pro Val Leu Ser Asp Asn Phe Thr Asp Ser Met Ile Met Asn Gly His Ser Pro Ile Ser Glu Gly Phe Gly Ala Arg Met Met 185 Ile Arg Phe Cys Pro Ser Cys Leu Asn Val Phe Asn Asn Val Gln Glu Asn Lys Asp Thr Ser Ile Phe Ser Arg Arg Ala Tyr Phe Ala Asp Pro Ala Leu Thr Leu Met His Glu Leu Ile His Val Leu His Gly Leu Tyr 230 235 Gly Ile Lys Ile Ser Asn Leu Pro Ile Thr Pro Asn Thr Lys Glu Phe 245 Phe Met Gln His Ser Asp Pro Val Gln Ala Glu Glu Leu Tyr Thr Phe Gly Gly His Asp Pro Ser Val Ile Ser Pro Ser Thr Asp Met Asn Ile 275 Tyr Asn Lys Ala Leu Gln Asn Phe Gln Asp Ile Ala Asn Arg Leu Asn Ile Val Ser Ser Ala Gln Gly Ser Gly Ile Asp Ile Ser Leu Tyr Lys Gln Ile Tyr Lys Asn Lys Tyr Asp Phe Val Glu Asp Pro Asn Gly Lys 330 Tyr Ser Val Asp Lys Asp Lys Phe Asp Lys Leu Tyr Lys Ala Leu Met Phe Gly Phe Thr Glu Thr Asn Leu Ala Gly Glu Tyr Gly Ile Lys Thr 355 Arg Tyr Ser Tyr Phe Ser Glu Tyr Leu Pro Pro Ile Lys Thr Glu Lys

Leu Leu Asp Asn Thr Ile Tyr Thr Gln Asn Glu Gly Phe Asn Ile Ala 390 Ser Lys Asn Leu Lys Thr Glu Phe Asn Gly Gln Asn Lys Ala Val Asn 410 Lys Glu Ala Tyr Glu Glu Ile Ser Leu Glu His Leu Val Ile Tyr Arg Ile Ala Met Cys Lys Pro Val Met Tyr Lys Asn Thr Gly Lys Ser Glu Gln Cys Ile Ile Val Asn Asn Glu Asp Leu Phe Phe Ile Ala Asn Lys 455 Asp Ser Phe Ser Lys Asp Leu Ala Lys Ala Glu Thr Ile Ala Tyr Asn Thr Gln Asn Asn Thr Ile Glu Asn Asn Phe Ser Ile Asp Gln Leu Ile 490 Leu Asp Asn Asp Leu Ser Ser Gly Ile Asp Leu Pro Asn Glu Asn Thr Glu Pro Phe Thr Asn Phe Asp Asp Ile Asp Ile Pro Val Tyr Ile Lys Gln Ser Ala Leu Lys Lys Ile Phe Val Asp Gly Asp Ser Leu Phe Glu Tyr Leu His Ala Gln Thr Phe Pro Ser Asn Ile Glu Asn Leu Gln Leu Thr Asn Ser Leu Asn Asp Ala Leu Arg Asn Asn Asn Lys Val Tyr Thr 565 570 Phe Phe Ser Thr Asn Leu Val Glu Lys Ala Asn Thr Val Val Gly Ala Ser Leu Phe Val Asn Trp Val Lys Gly Val Ile Asp Asp Phe Thr Ser Glu Ser Thr Gln Lys Ser Thr Ile Asp Lys Val Ser Asp Val Ser Ile 615 Ile Ile Pro Tyr Ile Gly Pro Ala Leu Asn Val Gly Asn Glu Thr Ala Lys Glu Asn Phe Lys Asn Ala Phe Glu Ile Gly Gly Ala Ala Ile Leu Met Glu Phe Ile Pro Glu Leu Ile Val Pro Ile Val Gly Phe Phe Thr 660 665 Leu Glu Ser Tyr Val Gly Asn Lys Gly His Ile Ile Met Thr Ile Ser Asn Ala Leu Lys Lys Arg Asp Gln Lys Trp Thr Asp Met Tyr Gly Leu 690 Ile Val Ser Gln Trp Leu Ser Thr Val Asn Thr Gln Phe Tyr Thr Ile 715

Lys Glu Arg Met Tyr Asn Ala Leu Asn Asn Gln Ser Gln Ala Ile Glu
725 730 735

Lys Ile Ile Glu Asp Gln Tyr Asn Arg Tyr Ser Glu Glu Asp Lys Met 740 745 750

Asn Ile Asn Ile Asp Phe Asn Asp Ile Asp Phe Lys Leu Asn Gln Ser
755 760 765

Ile Asn Leu Ala Ile Asn Asn Ile Asp Asp Phe Ile Asn Gln Cys Ser 770 775 780

Ile Ser Tyr Leu Met Asn Arg Met Ile Pro Leu Ala Val Lys Lys Leu 785 790 795 800

Lys Asp Phe Asp Asp Asn Leu Lys Arg Asp Leu Leu Glu Tyr Ile Asp 805 810 815

Thr Asn Glu Leu Tyr Leu Leu Asp Glu Val Asn Ile Leu Lys Ser Lys 820 830

Val Asn Arg His Leu Lys Asp Ser Ile Pro Phe Asp Leu Ser Leu Tyr 835 840 845

Thr Lys Asp Thr Ile Leu Ile Gln Val Phe Asn Asn Tyr Ile Ser Asn 850 860

Ile Ser Ser Asn Ala Ile Leu Ser Leu Ser Tyr Arg Gly Gly Arg Leu 865 870 875 880

Ile Asp Ser Ser Gly Tyr Gly Ala Thr Met Asn Val Gly Ser Asp Val 885 890 895

Ile Phe Asn Asp Ile Gly Asn Gly Gln Phe Lys Leu Asn Asn Ser Glu 900 905 910

Asn Ser Asn Ile Thr Ala His Gln Ser Lys Phe Val Val Tyr Asp Ser 915 920 925

Met Phe Asp Asn Phe Ser Ile Asn Phe Trp Val Arg Thr Pro Lys Tyr 930 940

Asn Asn Asn Asp Ile Gln Thr Tyr Leu Gln Asn Glu Tyr Thr Ile Ile 945 950 955 960

Ser Cys Ile Lys Asn Asp Ser Gly Trp Lys Val Ser Ile Lys Gly Asn 965 970 975

Arg Ile Ile Trp Thr Leu Ile Asp Val Asn Ala Lys Ser Lys Ser Ile 980 985 990

Phe Phe Glu Tyr Ser Ile Lys Asp Asn Ile Ser Asp Tyr Ile Asn Lys 995 1000 1005

Trp Phe Ser Ile Thr Ile Thr Asn Asp Arg Leu Gly Asn Ala Asn Ile 1010 1015 1020

Tyr Ile Asn Gly Ser Leu Lys Lys Ser Glu Lys Ile Leu Asn Leu Asp 1025 1030 1035 1040

Arg Ile Asn Ser Ser Asn Asp Ile Asp Phe Lys Leu Ile Asn Cys Thr 1045 1050 1055 Asp Thr Thr Lys Phe Val Trp Ile Lys Asp Phe Asn Ile Phe Gly Arg

Glu Leu Asn Ala Thr Glu Val Ser Ser Leu Tyr Trp Ile Gln Ser Ser 1075 1080 1085

Thr Asn Thr Leu Lys Asp Phe Trp Gly Asn Pro Leu Arg Tyr Asp Thr 1090 1095 1100

Gln Tyr Tyr Leu Phe Asn Gln Gly Met Gln Asn Ile Tyr Ile Lys Tyr 1105 1110 1115 1120

Phe Ser Lys Ala Ser Met Gly Glu Thr Ala Pro Arg Thr Asn Phe Asn 1125 1130 1135

Asn Ala Ala Ile Asn Tyr Gln Asn Leu Tyr Leu Gly Leu Arg Phe Ile 1140 1145 1150

Ile Lys Lys Ala Ser Asn Ser Arg Asn Ile Asn Asn Asp Asn Ile Val 1155 1160 1165

Arg Glu Gly Asp Tyr Ile Tyr Leu Asn Ile Asp Asn Ile Ser Asp Glu 1170 1180

Ser Tyr Arg Val Tyr Val Leu Val Asn Ser Lys Glu Ile Gln Thr Gln 1185 1190 1195 1200

Leu Phe Leu Ala Pro Ile Asn Asp Asp Pro Thr Phe Tyr Asp Val Leu 1205 1210 1215

Gln Ile Lys Lys Tyr Tyr Glu Lys Thr Thr Tyr Asn Cys Gln Ile Leu 1220 1225 1230

Cys Glu Lys Asp Thr Lys Thr Phe Gly Leu Phe Gly Ile Gly Lys Phe 1235 1240 1245

Val Lys Asp Tyr Gly Tyr Val Trp Asp Thr Tyr Asp Asn Tyr Phe Cys 1250 1255 1260

Ile Ser Gln Trp Tyr Leu Arg Arg Ile Ser Glu Asn Ile Asn Lys Leu 1265 1270 1275 1280

Arg Leu Gly Cys Asn Trp Gln Phe Ile Pro Val Asp Glu Gly Trp Thr 1285 1290 1295

Glu

(2) INFORMATION FOR SEQ ID NO:78:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 1535 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: double
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: DNA (genomic)
- (ix) FEATURE:
 - (A) NAME/KEY: CDS
 - (B) LOCATION: 108..1526

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:78:

AGA	TCTC	GAT	CCCG	CGAA	AT I	'AATA	CGAC	T CA	CTAT	'AGGG	GAA	TTGT	GAG	CGGA	TAACA	A	60
TTC	CCCT	'CTA	GAAA	TAAT	TT I	GTTT	'AACT	T TA	AGAA	.GGAG	ATA	TACC		Gly	CAT		116
CAT His	CAT His	His	CAT His	CAT His	CAT His	CAT His 10	His	CAC His	AGC Ser	AGC Ser	GGC Gly 15	' His	ATC Ile	GAA Glu	GGT Gly		164
CGT Arg 20	His	ATG Met	GCT Ala	AGC Ser	ATG Met 25	Ala	GAC Asp	ACA Thr	ATT	TTA Leu 30	Ile	CAA Gln	GTT Val	TTT Phe	AAT Asn 35		212
AAT Asn	TAT Tyr	ATT	AGT Ser	AAT Asn 40	ATT Ile	AGT Ser	AGT Ser	AAT Asn	GCT Ala 45	ATT Ile	TTA Leu	AGT Ser	TTA Leu	AGT Ser 50	TAT Tyr		260
AGA Arg	GGT Gly	GGG Gly	CGT Arg 55	TTA Leu	ATA Ile	GAT Asp	TCA Ser	TCT Ser 60	GGA Gly	TAT Tyr	GGT Gly	GCA Ala	ACT Thr 65	ATG Met	AAT Asn		308
GTA Val	GGT Gly	TCA Ser 70	GAT Asp	GTT Val	ATC Ile	TTT Phe	AAT Asn 75	GAT Asp	ATA Ile	GGA Gly	AAT Asn	GGT Gly 80	Gln	TTT Phe	AAA Lys		356
TTA Leu	AAT Asn 85	AAT Asn	TCT Ser	GAA Glu	AAT Asn	AGT Ser 90	AAT Asn	ATT Ile	ACG Thr	GCA Ala	CAT His 95	CAA Gln	AGT Ser	AAA Lys	TTC Phe		404
GTT Val 100	GTA Val	TAT Tyr	GAT Asp	AGT Ser	ATG Met 105	TTT Phe	GAT Asp	AAT Asn	TTT Phe	AGC Ser 110	ATT Ile	AAC Asn	TTT Phe	TGG Trp	GTA Val 115		452
AGG Arg	ACT Thr	CCT Pro	AAA Lys	TAT Tyr 120	AAT Asn	AAT Asn	AAT Asn	GAT Asp	ATA Ile 125	CAA Gln	ACT Thr	TAT Tyr	CTT Leu	CAA Gln 130	AAT Asn		500
GAG Glu	TAT Tyr	ACA Thr	ATA Ile 135	ATT Ile	AGT Ser	TGT Cys	ATA Ile	AAA Lys 140	AAT Asn	GAC Asp	TCA Ser	GGA Gly	TGG Trp 145	AAA Lys	GTA Val		548
TCT	ATT	AAG Lys 150	GGA Gly	AAT Asn	AGA Arg	ATA Ile	ATA Ile 155	TGG Trp	ACA Thr	TTA Leu	ATA Ile	GAT Asp 160	GTT Val	AAT Asn	GCA Ala		596
AAA Lys	TCT Ser 165	AAA Lys	TCA Ser	ATA Ile	TTT Phe	TTC Phe 170	GAA Glu	TAT Tyr	AGT Ser	ATA Ile	AAA Lys 175	GAT Asp	AAT Asn	ATA Ile	TCA Ser		644
GAT Asp 180	TAT Tyr	ATA Ile	AAT Asn	AAA Lys	TGG Trp 185	TTT Phe	TCC Ser	ATA Ile	ACT Thr	ATT Ile 190	ACT Thr	AAT Asn	GAT Asp	AGA Arg	TTA Leu 195		692
GGT Gly	AAC Asn	GCA Ala	AAT Asn	ATT Ile 200	TAT Tyr	ATA Ile	AAT Asn	GGA Gly	AGT Ser 205	TTG Leu	AAA Lys	AAA Lys	AGT Ser	GAA Glu 210	AAA Lys		740

AT Il	T TT e Le	A AA u As	C TT. n Let 21:	u AS	r AGA	A ATT	AAT Asr	TC' Se: 22	r Se	T AA' C Ası	T GA' n Ası	T AT	A GAG B Asp 225	Phe	C AAA e Lys	788
TT. Le	A AT	T AA e As 23	n cy:	r ACA	A GAT	T ACT	Thr 235	. TA:	A TTT	r GTT e Val	T TGO	3 ATT	Lys	GA: Asp	r TTT Phe	836
AA: Asi	T AT:		T GGT e Gly	r AGA ⁄ Arg	GAA Glu	TTA Leu 250	AST	GCT Ala	r ACA	GAA Glu	A GTA 1 Val 255	. Ser	TCA Ser	CTA Leu	TAT Tyr	884
TG(Tr ₁ 26(, 116	CA Gli	A TCA	TCT Ser	ACA Thr 265	Asn	ACT Thr	TTA	A AAA 1 Lys	GAT Asp 270) Phe	TGG Trp	GGG Gly	AAT Asn	CCT Pro 275	932
TTA Leu	AGA Arg	A TAC	C GAT	ACA Thr 280	GIN	TAC Tyr	TAT	CTG Leu	Phe	Asn	CAA Gln	GGT Gly	ATG Met	CAA Gln 290	AAT Asn	980
ATO	TAT Tyr	ATA Ile	A AAG Lys 295	Tyr	TTT Phe	AGT Ser	AAA Lys	GCT Ala 300	Ser	ATG Met	GGG Gly	GAA Glu	ACT Thr 305	GCA Ala	CCA Pro	1028
CGT Arg	ACA Thr	AAC Asn 310	Pne	AAT Asn	AAT Asn	GCA Ala	GCA Ala 315	ATA Ile	AAT Asn	TAT Tyr	CAA Gln	AAT Asn 320	TTA Leu	TAT	CTT Leu	1076
GGT Gly	TTA Leu 325	Arg	TTT Phe	ATT Ile	ATA Ile	AAA Lys 330	AAA Lys	GCA Ala	TCA Ser	AAT Asn	TCT Ser 335	CGG Arg	AAT Asn	ATA Ile	AAT Asn	1124
AAT Asn 340	мsр	AAT Asn	ATA Ile	GTC Val	AGA Arg 345	GAA Glu	GGA Gly	GAT Asp	TAT Tyr	ATA Ile 350	TAT Tyr	CTT Leu	AAT Asn	ATT Ile	GAT Asp 355	1172
AAT Asn	ATT Ile	TCT	GAT Asp	GAA Glu 360	TCT Ser	TAC Tyr	AGA Arg	GTA Val	TAT Tyr 365	GTT Val	TTG Leu	GTG Val	AAT Asn	TCT Ser 370	AAA Lys	1220
GAA Glu	ATT Ile	CAA Gln	ACT Thr 375	CAA Gln	TTA Leu	TTT Phe	Leu	GCA Ala 380	CCC Pro	ATA Ile	AAT Asn	GAT Asp	GAT Asp 385	CCT Pro	ACG Thr	1268
TTC Phe	TAT Tyr	GAT Asp 390	GTA Val	CTA Leu	CAA Gln	11e	AAA Lys 395	AAA Lys	TAT Tyr	TAT Tyr	GAA Glu	AAA Lys 400	ACA Thr	ACA Thr	TAT Tyr	1316
AAT Asn	TGT Cys 405	CAG Gln	ATA Ile	CTT Leu	Cys	GAA Glu 410	AAA (Lys)	GAT Asp	ACT Thr	Lys	ACA Thr 415	TTT Phe	GGG Gly	CTG Leu	TTT Phe	1364
GGA Gly 420	ATT Ile	GGT Gly	AAA Lys	Pne	GTT Val 425	AAA (Lys)	GAT '	ГАТ Гуг	GGA Gly	TAT Tyr 430	GTT Val	TGG Trp	GAT . Asp '	Thr	TAT Tyr 435	1412
GAT Asp	AAT Asn	TAT Tyr	Pne	TGC . Cys 440	ATA :	AGT (Ser (CAG 1	rp	TAT Tyr :	CTC . Leu .	AGA . Arg .	AGA . Arg	Ile :	CT Ser	GAA Glu	1460

.

AAT ATA AAA TTA AGG TTG GGA TGT AAT TGG CAA TTC ATT CCC GTG Asn Ile Asn Lys Leu Arg Leu Gly Cys Asn Trp Gln Phe Ile Pro Val 455 460 465

1508

GAT GAA GGA TGG ACA GAA TAACTCGAG Asp Glu Gly Trp Thr Glu 470

1535

(2) INFORMATION FOR SEQ ID NO:79:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 473 amino acids
 - (B) TYPE: amino acid
 - (D) TOPOLOGY: linear
- (ii) MOLECULE TYPE: protein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:79:

Met Gly His His His His His His His His His Ser Ser Gly His

1 10 15

Ile Glu Gly Arg His Met Ala Ser Met Ala Asp Thr Ile Leu Ile Gln
20 25 30

Val Phe Asn Asn Tyr Ile Ser Asn Ile Ser Ser Asn Ala Ile Leu Ser 35 40 45

Leu Ser Tyr Arg Gly Gly Arg Leu Ile Asp Ser Ser Gly Tyr Gly Ala
50 55 60

Thr Met Asn Val Gly Ser Asp Val Ile Phe Asn Asp Ile Gly Asn Gly 65 70 75 80

Gln Phe Lys Leu Asn Asn Ser Glu Asn Ser Asn Ile Thr Ala His Gln
85 90 95

Ser Lys Phe Val Val Tyr Asp Ser Met Phe Asp Asn Phe Ser Ile Asn 100 105 110

Phe Trp Val Arg Thr Pro Lys Tyr Asn Asn Asn Asp Ile Gln Thr Tyr 115 120 125

Leu Gln Asn Glu Tyr Thr Ile Ile Ser Cys Ile Lys Asn Asp Ser Gly
130 135 140

Trp Lys Val Ser Ile Lys Gly Asn Arg Ile Ile Trp Thr Leu Ile Asp 145 150 155 160

Val Asn Ala Lys Ser Lys Ser Ile Phe Phe Glu Tyr Ser Ile Lys Asp 165 170 175

Asn Ile Ser Asp Tyr Ile Asn Lys Trp Phe Ser Ile Thr Ile Thr Asn 180 185 190

Asp Arg Leu Gly Asn Ala Asn Ile Tyr Ile Asn Gly Ser Leu Lys Lys 195 200 205

Ser Glu Lys Ile Leu Asn Leu Asp Arg Ile Asn Ser Ser Asn Asp Ile 210 215 220 Asp Phe Lys Leu Ile Asn Cys Thr Asp Thr Thr Lys Phe Val Trp Ile 225 230 235

Lys Asp Phe Asn Ile Phe Gly Arg Glu Leu Asn Ala Thr Glu Val Ser 245 250 255

Ser Leu Tyr Trp Ile Gln Ser Ser Thr Asn Thr Leu Lys Asp Phe Trp
260 265 270

Gly Asn Pro Leu Arg Tyr Asp Thr Gln Tyr Tyr Leu Phe Asn Gln Gly
275 280 285

Met Gln Asn Ile Tyr Ile Lys Tyr Phe Ser Lys Ala Ser Met Gly Glu 290 295 300

Thr Ala Pro Arg Thr Asn Phe Asn Asn Ala Ala Ile Asn Tyr Gln Asn 305 310 315 320

Leu Tyr Leu Gly Leu Arg Phe Ile Ile Lys Lys Ala Ser Asn Ser Arg 325 330 335

Asn Ile Asn Asn Asp Asn Ile Val Arg Glu Gly Asp Tyr Ile Tyr Leu 340 345 350

Asn Ile Asp Asn Ile Ser Asp Glu Ser Tyr Arg Val Tyr Val Leu Val 355 360 365

Asn Ser Lys Glu Ile Gln Thr Gln Leu Phe Leu Ala Pro Ile Asn Asp 370 375 380

Asp Pro Thr Phe Tyr Asp Val Leu Gln Ile Lys Lys Tyr Tyr Glu Lys 385 390 395 400

Thr Thr Tyr Asn Cys Gln Ile Leu Cys Glu Lys Asp Thr Lys Thr Phe
405 410 415

Gly Leu Phe Gly Ile Gly Lys Phe Val Lys Asp Tyr Gly Tyr Val Trp
420 425 430

Asp Thr Tyr Asp Asn Tyr Phe Cys Ile Ser Gln Trp Tyr Leu Arg Arg
435 440 445

Ile Ser Glu Asn Ile Asn Lys Leu Arg Leu Gly Cys Asn Trp Gln Phe
450 455 460

Ile Pro Val Asp Glu Gly Trp Thr Glu
465 470

(2) INFORMATION FOR SEQ ID NO:80:

- (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 30 base pairs
 - (B) TYPE: nucleic acid
 - (C) STRANDEDNESS: single
 - (D) TOPOLOGY: linear

(ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "DNA"

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:80:

CGCCATGGCT GACACAATTT TAATACAAGT

- (2) INFORMATION FOR SEQ ID NO:81:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 32 base pairs (B) TYPE: nucleic acid (C) STRANDEDNESS: single

 - (D) TOPOLOGY: linear
 - (ii) MOLECULE TYPE: other nucleic acid (A) DESCRIPTION: /desc = "DNA"
 - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:81:

GCCTCGAGTT ATTCTGTCCA TCCTTCATCC AC

- (2) INFORMATION FOR SEQ ID NO:82:
 - (i) SEQUENCE CHARACTERISTICS:
 - (A) LENGTH: 12 amino acids
 - (B) TYPE: amino acid
 - (C) STRANDEDNESS: not relevant
 - (D) TOPOLOGY: not relevant
 - (ii) MOLECULE TYPE: peptide
 - (ix) FEATURE:
 - (A) NAME/KEY: Modified-site
 - (B) LOCATION: 12
 - (D) OTHER INFORMATION: /note= "The asparagine residue at this position contains an amide group."

32

(xi) SEQUENCE DESCRIPTION: SEQ ID NO:82:

Cys Gln Thr Ile Asp Gly Lys Lys Tyr Tyr Phe Asn